dihydrate crystal deposition disease AUTHOR(S):

Ryan, Lawrence M.; Lynch, Michael P.;

McCarty, Daniel J.

CORPORATE SOURCE: Dep. Med., Med. Coll. Wisconsin, Milwaukee, WI, USA

SOURCE: Arthritis Rheum. (1983), 26(4), 564-6

CODEN: ARHEAW; ISSN: 0004-3591

DOCUMENT TYPE: Journal LANGUAGE: English

CLASSIFICATION: 14-11 (Mammalian Pathological Biochemistry)

Section cross-reference(s): 13

ABSTRACT:

The total intracellular inorg, pyrophosphate (PPi) content and the fraction of PPi released (by thrombin) from the storage pool of blood platelets from normal humans, patients with sporadic Ca pyrophosphate dihydrate crystal deposition, and patients with familial disease were studied. No differences in group ranges or means were found. There appeared to be a rise in platelet PPi content with age of normal humans.

SUPPL. TERM: blood platelet inorg pyrophosphate; calcium pyrophosphate

deposition disease platelet

INDEX TERM: Senescence and Semility

(inorg. pyrophosphate of blood platelet in numans in

relation to;

INDEX TERM: Pseudogout

(inorg. pyrophosphate of blood platelet in, in humans)

INDEX TERM: Blood platelet

(inorg. pyrophosphate of, in primary and secondary

chondrocalcinosis and health in human)

INDEX TERM: 17031-92-4

ROLE: BIOL (Biological study)

(metabolic disorders, crystal deposition disease, inorq.

pyrophosphate of blood platelets in, in human,

14000-31-8 INDEX TERM:

ROLE: BIOL (Biological study)

(of blood platelets, in familial and secondary

chondrocalcinosis and health in human)

ANSWER 7 OF 28 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: 1998:414091 BIOSIS DOCUMENT NUMBER:

PREV199800414091

TITLE:

Scanning probe microscopy in biotechnology.

AUTHOR(S):

Henderson, Eric (1)

CORPORATE SOURCE:

(1) Iowa State Univ., Ames, IA USA

SOURCE:

Scanning, (April, 1998) Vol. 20, No. 3, pp. 138. Meeting Info.: Scanning 98: Meeting of the Foundation for Advances in Medicine and Science Baltimore, Maryland, USA May 10-12, 1998 Foundation for Advances in Medicine and

Science

. ISSN: 0161-0457.

DOCUMENT TYPE:

Conference English

LANGUAGE: CONCEPT CODE:

Microscopy Techniques - General and Special Techniques

\*01052

Genetics and Cytogenetics - General \*03502 Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biophysics - General Biophysical Techniques \*10504

Biophysics - Bioengineering \*10511

General Biology - Symposia, Transactions and Proceedings of

Conferences, Congresses, Review Annuals \*00520

INDEX TERMS:

Major Concepts

Bioprocess Engineering; Methods and Techniques

INDEX TERMS: Methods & Equipment

atomic force microscopy: microscopy method

INDEX TERMS:

Miscellaneous Descriptors

biotechnology; Meeting Abstract

L2 ANSWER 17 OF 28 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: 1997:119181 BIOSIS DOCUMENT NUMBER: PREV199799425684

TITLE: Chicken erythrocyte nucleosomes have a defined orientation

along the linker DNA: A scanning force microscopy study.

AUTHOR(S): Fritzche, Wolfgang (1); Henderson, Eric

CORPORATE SOURCE: (1) Inst. Phys. High Technol., Dep. Cryoelectronics

Microsystems, P.O. Box 100 239, D-07702 Jena Germany

SOURCE: Scanning, (1997) Vol. 19, No. 1, pp. 42-47.

ISSN: 0161-0457.

DOCUMENT TYPE:

Article English

LANGUAGE: ABSTRACT:

The orientation of nucleosomes was investigated using scanning force microscopy (SFM) of hypotonically spread chicken chromatin. A virtual cross section parallel to the substrate at half maximum height of the nucleosomal structure revealed an elliptical shape. The orientation of the major axis of this ellipse was investigated in reference to the direction of the axis of the nucleosoma, chain. An alignment of the nucleosomes along the nucleosomal chain was observed, with more than 50% of the nucleosomes aligned with the long axis of the chain within ltoreq 30 degree deviation. The alignment distribution peaked at 10-20 degree. The application of SFM-based image processing for the structural investigation of a protein-DNA complex demonstrates the potential

for this approach in structural molecular biology.

CONCEPT CODE: Mic

Microscopy Techniques - General and Special Techniques

\*01052

Cytology and Cytochemistry - Animal \*02506 Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biochemical Studies - Proteins, Peptides and Amino Acids

\*10064

Blood, Blood-Forming Organs and Body Fluids - Blood Cell

Studies \*15004

BIOSYSTEMATIC CODE: Galliformes \*85536

INDEX TERMS:

Major Concepts

Biochemistry and Molecular Biophysics; Blood and Lymphatics

(Transport and Circulation); Cell Biology; Methods and

Techniques

INDEX TERMS: Miscellaneous Descriptors

BLOOD AND LYMPHATICS; CELL BIOLOGY; CHROMATIN; DNA; DNA-PROTEIN COMPLEX; ERYTHROCYTE; IMAGE PROCESSING;

METHODOLOGY; MICROSCOPY METHOD; NUCLEOSOMES; SCANNING FORCE

MICROSCOPY

ORGANISM: Super Taxa

Galliformes: Aves, Vertebrata, Chordata, Animalia

ORGANISM:

Organism Name

chicken (Galliformes)

ORGANISM: Organism Superterms

animals; birds; chordates; nonhuman vertebrates;

vertebrates

ANSWER 22 OF 28 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: 1995:315199 BIOSIS DOCUMENT NUMBER: PREV199598329499

TITLE: Imaging and manipulating chromosomes with the atomic force

microscope.

AUTHOR(S): Jondle, Daniel M.; Ambrosio, Linda; Vesenka, James;

Henderson, Eric (1)

(1) Dep. Zool. Genet., Iowa State Univ., Ames, IA 50011 USA Chromosome Research, 11995; Vol. 3, No. 4, pp. 239-244. CORPORATE SOURCE: SOURCE:

ISSN: 0967-3849.

DOCUMENT TYPE:

Article English

LANGUAGE: ABSTRACT:

Polytene chromosomes from the salivary gland cells of Drosophila melanogaster were examined by atomic force microscopy. The atomic force microscope (AFM) was capable of resolving chromosomal features down to the limits of the tip sharpness, about 500 ANG for pyramidal-shaped tips. Resolution was increased to 300 ANG by using electron beam deposited (EBD) tips with high aspect ratios. This significantly exceeds the resolution obtainable with conventional optical microscopes, but at the cost of compromising the structural integrity of the sample. A reasonable compromise was achieved by using exide-sharpened tips. In this case high resolution was obtained without sample degradation, but when desired these tips were also capable of sample disintegration with increased scanning force and rate. Thus, oxide-sharpened tips were used to precisely dissect defined chromosomal regions to illustrate their potential use in genetic mapping efforts. This study illustrates the utility of the AFM in the characterization and manipulation of chromosomes and chromosomal DNA. CONCEPT CODE:

Microscopy Techniques - General and Special Techniques

01052

Genetics and Cytogenetics - Animal \*03506 Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines 10062

Anatomy and Histology, General and Comparative -Microscopic and Ultramicroscopic Anatomy \*11108

Invertebrata, Comparative and Experimental Morphology, Physiology and Pathology - Insecta - Physiology \*64076

BIOSYSTEMATIC CODE: Diptera \*75314

INDEX TERMS: Major Concepts

Genetics; Morphology; Physiology

Miscellaneous Descriptors INDEX TERMS:

POLYTENE CHROMOSOME; ULTRASTRUCTURE

ORGANISM: Super Taxa

Diptera: Insecta, Arthropoda, Invertebrata, Animalia

ORGANISM: Organism Name

Drosophila melanogaster (Diptera)

ORGANISM: Organism Superterms

animals; arthropods; insects; invertebrates

L3 ANSWER 1 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: 2000:544893 BIOSIS DOCUMENT NUMBER: PREV200000544893

TITLE: The effect of super-oxidized water on Escherichia coli.

AUTHOR(S): Zinkevich, V. (1); Beech, I. B.; Tapper, R.; Bogdarina, I.

CORPORATE SCURCE: (1) School of Pharmacy and Biomedical Sciences, University

of Portsmouth, White Swan Road, St. Michael's Building,

Portsmouth, PO1 2DT UK

SOURCE: Journal of Hospital Infection, (October, 2000) Vol. 46, No.

2, pp. 153-156. print.

ISSN: 0195-6701.

DOCUMENT TYPE: Article LANGUAGE: English SUMMARY LANGUAGE: English

ABSTRACT:

The mechanism of action of Sterilox, a non-toxic liquid blocide produced by electrolysis of a dilute saline solution, upon planktonic cells of Escherichia coli JM109 was investigated using protein and nucleic acid analysis.

The results revealed total destruction of chromosomal and plasmid DNA

, RNA and proteins of E. coli within 5 min of exposure. Our earlier

investigation conducted using atomic force

\*\*\*microscopy\*\*\* imaging revealed swelling and rupture of E. coli cells with release of cytoplasm. We propose that the biocidal properties of Sterilox are due to its effect upon constituents of the bacterial cell including proteins and nucleic acids.

CONCEPT CODE: Physiology and Biochemistry of Bacteria \*31000

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biochemical Studies - Proteins, Peptides and Amino Acids

\*10064

Public Health: Environmental Health - Sewage Disposal and

Sanitary Measures \*37014

Pest Control, General; Pesticides; Herbicides \*54600

BIOSYSTEMATIC CODE: Enterobacteriaceae 06702

INDEX TERMS: Major Concepts

Sanitation

INDEX TERMS: Chemicals & Biochemicals

Sterilox: biocide; plasmid DNA; plasmid RNA;

proteins; super-oxidized water

INDEX TERMS: Methods & Equipment

electrolysis: synthetic method; nucleic acid

analysis: analytical method; protein analysis: analytical

method

ORGANISM: Super Taxa

Enterobacteriaceae: Facultatively Anaerobic Gram-Negative Rods, Eubacteria, Bacteria, Microorganisms; Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia

ORGANISM: Organism Name

Escherichia coli (Enterobacteriaceae): pathogen,

strain-JM109; human (Hominidae): patient

ORGANISM: Organism Superterms

Animals; Bacteria; Chordates; Eubacteria; Humans; Manmala;

Microorganisms; Primates; Vertebrates

L3 ANSWER 2 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: 2000:497584 BIOSIS
DOCUMENT NUMBER: PREV200000497705

TITLE: Near field microscopies: From isolated molecules to livin:

cells.

AUTHOR(S): Delain, E. (1); Michel, D. (1); Le Grimellec, Ch. CORPORATE SOURCE: (1) Laboratoire de Microscopie Moleculaire et

Cellulaire/CNRS UMR 8532, Institut Gustave Roussy, Rue

Camille Desmoulins, Villejuif France

SOURCE: Morphologie, (Juin, 2000) Vol. 84, No. 265, pp. 25-30.

print.

ISSN: 1286-0115.

DOCUMENT TYPE: LANGUAGE:

Article French

SUMMARY LANGUAGE: English; French

ABSTRACT:

Near field (or scanning probe) microscopy is a recent technology which, owing to the huge amount of publications, is becoming a reference method in molecular and cellular imaging. These microscopies consist in the scanning of the sample, line by line, with a very tiny tip and thus providing informations on its surface down to the nanometer scale. These methods gather scanning tunelling microscopy (STM), which measures a current between the tip and the specimen support, atomic force (AFM), which measures the repulsive and attractive forces of \*\*\*microscopy\*\*\* the tip in contact or very close to the specimen, and scanning near field optical microscopies (SNOM), for which a glass tip allows to catch light signals. Atomic force microscopy, which allows the observation of specimens in air or physiological conditions environments,

is presently dominant in biology, in complementarity with the classical optical and electron microscopies, which by the way, have also shown considerable improvements during the last years. The complementarity of these microscopie: is due to their very different basic principles, which provide them various possibilities and limits. The biological applications of STM is limited by the need of conducting samples, but the different models of SNOM, often still is development, allow to consider very interesting applications, particularly for detecting very faint and tiny fluorescence signals. Different examples will kn given concerning the visualization by AFM of isolated DNA molecules, naked or associated with proteins, the observation of intact or decondensed chromosomes, as well as living cells. One of the originality of AFM is its capacity to observed objects in a wide range of enlargements, with fields from a few hundred of nanometers to several micrometers.

CONCEPT CODE:

Cytology and Cytochemistry - General

Biochemical Studies - General \*10060

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biochemical Studies - Proteins, Peptides and Amino Acids

\*10C64

INDEX TERMS: Major Concepts

Biochemistry and Molecular Biophysics; Cell Biology;

Methods and Techniques

INDEX TERMS: Parts, Structures, & Systems of Organisms

cells: near field microscopic study

INDEX TERMS:

Chemicals & Biochemicals

double stranded DNA: microscopic study; nucleic acids: microscopic study; proteins:

microscopic study

INDEX TERMS: Methods & Equipment

scanning near field optical microscopy:

analytical method; scanning probe microscopy (near field microscopy): analytical method

ORGANISM:

Super Taxa Organisms

ORGANISM:

Organism Name

organism (Organisms)

ANSWER 3 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS L3

ACCESSION NUMBER: DOCUMENT NUMBER:

2000:309262 BIOSIS PREV200000309262

TITLE:

A dimer as a building block in assembling RNA: A hexamer

that gears bacterial virus phi29 DNA

-translocating machinery.

AUTHOR(S):

SOURCE:

Chen, Chaoping; Sheng, Sitong; Shao, Zhifeng; Gut, Peixust. Journal of Biological Chemistry, (June 9, 2006, Vol. 2/5,

No. 23, pp. 17510-17516. print.

ISSN: 0021-9258.

DOCUMENT TYPE:

Article English English

LANGUAGE: SUMMARY LANGUAGE:

ABSTRACT:

Six RNA (pRNA) molecules form a hexamer, via hand-in-hand interaction, to gear bacterial virus phi29 DNA translocation machinery. Here we report the pathway and the conditions for the hexamer formation. Stable pRNA dimers and trimers were assembled in solution, isolated from native gels, and separated L, sedimentation, providing a model system for the study of RNA dimers and trimers in a protein-free environment. Cryoatomic force microscopy revealed that monomers displayed a check-mark shaped outline, dimers exhibited an elongated shape, and trimers formed a triangle. Dimerization of pRNA was promoted by a variety of cations including spermidine, whereas procapsid binding and DNA packaging required specific divalent cations, including Mg2+, Ca2+, and Mn2+. Both the tandem and fused pRNA dimers with complementary loops designed to form even-numbered rings were active in packaging, whereas those without complementary loops were inactive. We conclude that dimers are the building blocks of the hexamer, and the pathway of building a hexamer is: dimer fwdarw tetramer fwdarw hexamer. The Hill coefficient of 2.5 suggests that there are three binding sites wit: 300perative binding on the surface of the procapsid. The two interacting loops played a key

foundation for hexamer assembly. CONCEPT CODE:

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Methods, Materials and Apparatus, General - Laboratory

Methods \*01004

Virology - Animal Host Viruses \*33506 Biochemical Studies - Minerals \*10069

role in recruiting the incoming dimer, whereas the procapsid served as the

INDEX TERMS:

Major Concepts

Biochemistry and Molecular Biophysics; Methods and

Techniques

INDEX TERMS:

Chemicals & Biochemicals

DNA-translocating machinery; RNA: assay,

isolation, separation, synthesis; calcium (LF); magness and

(II); manganese (II)

INDEX TERMS:

Methods & Equipment

PAGE [polyacrylamide gel electrophoresis]: gel

electrophoresis, isolation method; RNA binding assay: analytical method, binding assays; RNA synthesis: nucleic acid synthesis, synthetic method; cryo-

atomic force microscopy

[cryo-AFM]: microscopy method, microscopy : CB; dimer binding competition assay:

Analysis/Characterization Techniques: CB, analytical method; sucrose gradient sedimentation: Extraction, Isolation, Purification and Separation Techniques,

separation method; virion assembly assay:

Analysis/Characterization Techniques: CB, analytical method

ORGANISM: Super Taxa

Podoviridae: Bacterial Viruses, Viruses, Microorganisms

ORGANISM: Organism Name

> phi29 (Podoviridae) Organism Superterms

Bacterial Viruses; Microorganisms; Viruses

REGISTRY NUMBER:

ORGANISM:

14127-61-8 (CALCIUM (II)) 22537-22-0 (MAGNESIUM (II)) 16397-91-4 (MANGANESE (II))

ANSWER 4 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS L3

ACCESSION NUMBER: DOCUMENT NUMBER:

2000:246496 BIOSIS PREV200000246496

TITLE:

PNA-dependent gene chemistry: Stable coupling of peptide:

and oligonucleotides to plasmid DNA.

AUTHOR(S): Zelphati, O.; Liang, X.; Nguyen, C.; Barlow, S.; Sheng, S.;

Shao, Z.; Felgner, P. L. (1)

CORPORATE SOURCE: (1) Gene Therapy Systems, 10190 Telesis Court, San Diego,

CA, 92121 USA

SOURCE: Biotechniques, (Feb., 2000) Vol. 28, No. 2, pp. 304-316.

ISSN: 0736-6205.

DOCUMENT TYPE: A
LANGUAGE: E

SUMMARY LANGUAGE:

Article English English

ABSTRACT:

Two approaches are described for stably conjugating peptides, proteins and

oligonucleotides onto plasmid DNA. Both methods use a peptide

\*\*\*nucleic\*\*\* acid (PNA) clamp, which binds irreversibly and specifically to

a binding site cloned into the plasmid. The first approach uses a

biotin-conjugated PNA clamp that can be used to introduce functional biotin

groups onto the plasmid to which streptavidin can bind. Atomic

\*\*\*force\*\*\* microscopy images of linearized plasmid show streptavidin localized at the predicted PNA binding site on the DNA strand. Peptides and oligonucleotides containing free thiol groups were conjugated to maleimide streptavidin, and these streptavidin conjugates were bound to the biotin-PNA-labeled plasmid. In this way, peptides and oligonucleotides could be brought into stable association with the plasmid. A second approach used a maleimide-conjugated PNA clamp. Methods are described for conjugating thiolated peptides and oligonucleotides directly to the maleimide-PNA-DNA hybrid. This straightforward technology offers an easy approach to introduce functional groups onto plasmid DNA without

disturbing its transcriptional activity.

CONCEPT CODE: Biochemical Methods - Nucleic Acids, Purines and

Pyrimidines \*10052

Biochemical Studies - Proteins, Peptides and Amino Acids

\*10064

Biophysics - General Biophysical Techniques \*10504

Genetics of Bacteria and Viruses \*31500

INDEX TERMS: Major Concepts

Molecular Genetics (Biochemistry and Mclecular Biophysic: ;

Methods and Techniques

INDEX TERMS: Chemicals & Biochemicals

oligonucleotides; peptide nucleic acid; peptides;

plasmid DNA

INDEX TERMS: Methods & Equipment

ATTO-TAG labeling kit: Molecular Probes, equipment; agards-

gel electrophoresis: analytical method, gei

electrophoresis; atomic force microscopy: microscopy method, microscopy: CB, microscopy: CT;

transfection: gene expression/vector techniques, genetic

method; transmission electron microscopy:

electron microscopy: CB, electron microscopy: CT, microscopy method

INDEX TERMS:

Miscellaneous Descriptors PNA-dependent gene chemistry

L3 ANSWER 5 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: DOCUMENT NUMBER:

2000:61886 BIOSIS PREV200000061886

TITLE:

Polymerase activities and RNA structures in the

atomic force microscope.

AUTHOR(S):

Hansma, Helen G. (1); Golan, Roxana; Hsieh, Wan;

Daubendiek, Sarah L.; Kool, Eric T.

CORPORATE SOURCE:

(1) Department of Physics, University of California, Santa

Barbara, Santa Barbara, CA USA

SOURCE:

Journal of Structural Biology, (Oct., 1999, V.). 127, No.

3, pp. 240-247.

ISSN: 1047-8477.

DOCUMENT TYPE:

Article English

LANGUAGE: SUMMARY LANGUAGE:

English

ABSTRACT:

The structures of the reaction products are the basis for novel polymerase

assays using the atomic force microscope (AFM). Polymerases

are the enzymes involved in transcription and replication of DNA. Rapid semiquantitative estimates of the activity of DNA polymerases

such as Sequenase, Taq polymerase, and AMV reverse transcriptase and RNA

polymerases (RNAP) such as Escherichia coli RNAP were obtained from AFM images

of the nucleic acids after polymerase reactions. DNA

polymerases were assayed via replication of the single-stranded PHIX-174 virion. RNAP was assayed via transcription, using a rolling sirily DNA

template that produces long strands of RNA. In some cases, AFM was better than agarose gel electrophoresis for assaying DNA polymerase activity,

since aggregation prevented the DNA from entering the agarose ge..

Extended molecules of single-stranded RNA synthesized with the rolling circle template showed varied conformations and degrees of stretching. \* \* \* DNA \* \* \*

Some structural differences were observed between two RNAs-a ribozyme concatamer and an RNA with 90% purines.

CONCEPT CODE:

Genetics and Cytogenetics - General \*03502

Microscopy Techniques - Electron Microscopy \*01058 Biochemical Methods - Nucleic Acids, Purines and

Pyrimidines \*10052

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Replication, Transcription, Translation 18300 Biophysics - Molecular Properties and Macromolecules

\*10506

Enzymes - Methods \*18884

Enzymes - Chemical and Physical \*10806

Physiology and Biochemistry of Bacteria \*31000

Genetics of Bacteria and Viruses \*31500 Virology - General; Methods \*33502

BIOSYSTEMATIC CODE: Microviridae

02706

06702 Enterobacteriaceae

INDEX TERMS:

Major Concepts

Enzymology (Biochemistry and Molecular Biophysics);

Molecular Genetics (Biochemistry and Molecular Biophysics)

INDEX TERMS:

Chemicals & Biochemicals

AMV reverse transcriptase; DNA: rolling circle template; DNA polymerase; RNA: structure; RNA

polymerase; Taq polymerase

INDEX TERMS:

Methods & Equipment

atomic force microscope: laboratory

equipment; atomic force

microscopy: analytical method, microscopy

: CB, molecular imaging method

ORGANISM:

Super Taxa

Enterobacteriaceae: Facultatively Anaerobic Gram-Negative Rods, Eubacteria, Bacteria, Microorganisms; Microviridae:

Bacterial Viruses, Viruses, Microorganisms

ORGANISM:

Organism Name

Escherichia coli (Enterphanteria seae ; bantes: grave

phi-X-174 (Microviridae)

ORGANISM:

Organism Superterms

Bacteria; Bacterial Wirdses; Edbacteria; M. H. Garaganisto;

Viruses

REGISTRY NUMBER:

9012-90-2 (DNA POLYMERASE)

9014-24-8 (RNA POLYMERASE)

ANSWER 6 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: 2000:2359 BIOSIS DOCUMENT NUMBER:

PREV200000002359

TITLE:

DNA toroids: Stages in condensation.

AUTHOR(S):

Golan, Roxana; Pietrasanta, Lia I.; Hsieh, Wan; Hansma,

Helen G. (1)

CORPORATE SOURCE:

(1) Department of Physics, UCSB, Santa Barbara, CA, 93106

USA

SOURCE:

Biochemistry, (Oct. 19, 1999) Vol. 38, No. 42, pp.

14069-14076.

ISSN: 0006-2960.

DOCUMENT TYPE:

Article English

LANGUAGE:

SUMMARY LANGUAGE:

English

The effects of polylysine (PLL) and PLL-asialoorosomucoid (AsOR) on DNA condensation have been analyzed by AFM. Different types of condensed structures were observed, which show a sequence of conformational

changes as circular plasmid DNA molecules condense progressively. The structures range from circular molecules with the length of the plasmid to small toroids and short rods with apprx1/6 to 1/8 the contour

length of the uncondensed circular DNA. Single plasmid molecules of 6800 base pairs (bp) condense into single toroids of apprx110 nm diameter,

measured center-to-center. The results are consistent with a model for condensation in which circular DNA molecules fold several

times into progressively shorter rods. Structures intermediate between torolar and rods suggest that at least some toroids may form by the opening up of rods as proposed by Dunlap et al. ((1997) Nucleic Acids Res. 25, 3095).

Toroids and rods formed at lysine: nucleotide ratios of 5:1 and 6:1. This high lys ine: nucleotide ratio is discussed in relation to entropic considerations

and the overcharging of macroions. PLL-AsOR is much more effective than PLL alone for condensing DNA, because several PLL molecules are attached

to a single AsOR molecule, resulting in an increased cation density. CONCEPT CODE:

Genetics and Cytogenetics - General \*03502

Microscopy Techniques - General and Special Techniques

Biochemical Methods - General \*10050 Biochemical Studies - General \*10060

Biophysics - General Biophysical Studies \*10502

INDEX TERMS:

Major Concepts

Molecular Genetics (Biochemistry and Molecular Biophysics;

Methods and Techniques

Chemicals & Biochemicals INDEX TERMS:

DNA toroids: analysis, condensation stages;

polylysine: DNA condenser; polylysine-

asialoorosomucoid: DNA condenser

INDEX TERMS:

Methods & Equipment Fast Flow Q Sepharose anion exchange chromatography:

chromatographic techniques, separation method;

atomic force microscopy:

microscopy method, microscopy: CB

REGISTRY NUMBER:

25104-18-1Q (POLYLYSINE) 38000-06-5Q (POLYLYSINE)

ANSWER 7 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: DOCUMENT NUMBER:

1999:187181 BIOSIS PREV199900187181

TITLE:

Activity of a single gmoniclease revealed by atomic

force microscopy.

AUTHOR(S):

Takeuchi, M. (1); Okada, T. (1)

CORPORATE SOURCE:

(1) Joint Research Center for Atom Technology 3/6 MAIR,

1-1-4 Higashi, Tsukuba, Ibaraki, 305-0046 Japan -- --

SOURCE:

Biophysical Journal, (Jan., 1999) Vol. 76, No. 1 PART 2,

pp. A132.

Meeting Info.: Forty-third Annual Meeting of the Biophysical Society Baltimore, Maryland, USA February 13-17, 1999

ISSN: 0006-3495.

DOCUMENT TYPE:

Conference

LANGUAGE:

English

CONCEPT CODE:

Biochemical Studies - General \*10060 Biochemical Methods - General \*10050

Enzymes - General and Comparative Studies; Coenzymes

\*10802

General Biology - Symposia, Transactions and Proceedings of

Conferences, Congresses, Review Annuals \*00520

INDEX TERMS:

Major Concepts

Enzymology (Biochemistry and Molecular Biophysics); Methods

and Techniques

INDEX TERMS:

Chemicals & Biochemicals exonuclease: activity; DNA

INDEX TERMS:

Methods & Equipment

atomic force microscopy:
microscopy method, microscopy: CB

INDEX TERMS:

Miscellaneous Descriptors

enzyme kinetics; protein-nucleic acid

interactions; Meeting Abstract; Meeting Poster

REGISTRY NUMBER:

37228-74-3 (EXONUCLEASE)

L3 ANSWER 8 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: DOCUMENT NUMBER:

1998:256680 BICSIS PREV199800256680

TITLE:

Novel vectors for gene delivery formed by self-assembly of

DNA with poly(L-lysine) grafted with hydrophilic

polymers.

AUTHOR(S):

Toncheva, Veska; Wolfert, Margreet A.; Dash, Philip R.; Oupicky, David; Ulbrich, Karel; Seymour, Leonard W. (1);

Schacht, Etienne H.

CORPORATE SOURCE:

(1) CRC Inst. Cancer Studies, Univ. Birmingham, Birmingham

B15 2TA UK

SOURCE:

Biochimica et Biophysica Acta, (May 8, 1998) Vol. 138, No.

3, pp. 354-368. ISSN: 0006-3002.

DOCUMENT TYPE:

Article English

LANGUAGE: ABSTRACT:

Complexes formed between **DNA** and cationic polymers are attracting increasing attention as novel synthetic vectors for delivery of genes. We are trying to improve biological properties of such complexes by oriented self-assembly of **DNA** with cationic-hydrophilic block copolymers, designed to enshroud the complex within a protective hydrophilic polymer corona. Poly(L-lysine) (pLL) grafted with range of hydrophilic polymer blocks, including poly(ethylene glycol) (pEG), dextran and poly(N-(2-hydroxypropyl)methacrylamide) (pHPMA), shows efficient binding to **DNA** and mediates particle self-assembly and inhibition of ethidium bromide/\*\*\*DNA\*\*\* fluorescence. The complexes formed are discrete and typically about 100 nm diameter, viewed by **atomic force microscopy** 

. Surface charges are slightly shielded by the presence of the hydrophilic polymer, and complexes generally show decreased cytotoxicity compared with simple pLL/DNA complexes. pEG-containing complexes show increased transfection activity against cells in vitro. Complexes from a conjugates showed greater aqueous solubility than simple pLL/DNA complexes, particularly at charge neutrality. These materials appear to have the ability to regulate the physicochemical and biological properties of polycation/DNA complexes, and should find important applications in packaging of nucleic acids for specific biological applications.

CONCEPT CODE: Genetics and Cytogenetics - General \*03502

Microscopy Techniques - Electron Microscopy \*01058 Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biochemical Studies - General \*10060

INDEX TERMS: Major Concepts

Biochemistry and Molecular Biophysics

INDEX TERMS: Chemicals & Biochemicals

hydrophilic polymers; poly(L-lysine); DNA:

self-assembly

INDEX TERMS: Methods & Equipment

atomic force microscopy:

analytical method

INDEX TERMS: Miscellaneous Descriptors

gene delivery: novel vectors
25104-18-1Q (POLY(L-LYSINE))

REGISTRY NUMBER: 25104-18-1Q (POLY(L-LYSINE)) 38000-06-5Q (POLY(L-LYSINE))

L3 ANSWER 9 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER:

1998:256659 BIOSIS

DOCUMENT NUMBER:

PREV199800256659

TITLE:

Analysis of various sequence-specific triplexes by electron

and atomic force microscopies.

AUTHOR(S):

Cherny, Dimitry I. (1); Fourcade, Alain; Svinarchuk, Fedro;

Nielsen, Peter E.; Malvy, Claude; Delain, Etienne

CORPORATE SOURCE:

(1) Lab. Microscopie Cellularie Molecularie, URA 147, CNRS,

Inst. Gustave-Roussy, rue Camillie Descoulin, F- 440

Villejuif France

SOURCE:

Biophysical Journal, (Feb., 1998) Vol. 74, No. 2 PART 1,

pp. 1015-1023. ISSN: 0006-3495.

DOCUMENT TYPE:

Article

LANGUAGE:

English

ABSTRACT:

Sequence-specific interactions of 20-mer G, A-containing triple helix-forming oligonucleotides (TFOs) and bis-PNAs (peptide nucleic acids) with

double-stranded DNA was visualized by electron (EM) and

\*\*\*atomic\*\*\* force (AFM) microscopies. Triplexes formed by

biotinylated TFOs are easily detected by both EM and AFM in which streptavidin is a marker. AFM images of the unlabeled triplex within a long plasmid

\*\*\*DNA\*\*\* show a apprx0.4-nm height increment of the double helix within the target site position. TFOs conjugated to a 74-nt-long oligonucleotide forming a 33-bp-long hairpin form extremely stable triplexes with the target site that are readily imaged by both EM and AFM as protruding DNA. The short

duplex protrudes in a perpendicular direction relative to the double helix axis, either in the plane of the support or out of it. In the latter case, the apparent height of the protrusion is apprx 1.5 nm, when that of the crimlex site is increased by 0.3-0.4 nm. Triplex togration by pis-FVA, decamers of PNA are connected via a flexible linker, causes deformations of the

double helix at the target site, which is readily detected as kinks by both FM and AFM. Moreover, AFM shows that these kinks are often accompanied by an increase in the DNA apparent height of apprx 35%. This work shows the

first direct visualization of sequence-specific interaction of TFOs and PNAs, with their target sequences within long plasmid DNAs, through the measurements of the apparent height of the **DNA** double helix by AFM.

CONCEPT CODE:

Genetics and Cytogenetics - General \*03502

Microscopy Techniques - Electron Microscopy \*01058 Biochemical Methods - Nucleic Acids, Purines and

Pyrimidines \*10052

Biochemical Methods - Proteins, Peptides and Amino Acids

\*10054

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biochemical Studies - Proteins, Peptides and Amino Acids

\*10064

Biophysics - Molecular Properties and Macromolecules

\*10506

INDEX TERMS:

Major Concepts

Methods and Techniques; Molecular Genetics (Biochemistry

and Molecular Biophysics)
Chemicals & Biochemicals

bis-peptide nucleic acid: DNA

interaction; double-stranded DNA; triple
helix-forming oligonucleotide: DNA interaction

INDEX TERMS: Methods & Equipment

atomic force microscopy:

analytical method; electron microscopy:

analytical method

L3 ANSWER 10 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: DOCUMENT NUMBER:

1998:221880 BIOSIS PREV199800221880

TITLE:

Study of the interaction of **DNA** with displatinand other Pd(II) and Pt(II) complexes by **atomic** 

force microscopy.

AUTHOR(S):

INDEX TERMS:

Onoa, G. Bibiana; Cervantes, Gemma; Moreno, Virtudes (1);

Prieto, M. Jose

CORPORATE SOURCE:

(1) Dep. Quim. Inorg., Univ. Barcelona, Diagonal 647,

08028-Barcelona Spain

SOURCE:

Nucleic Acids Research, (March 15, 1998) Vol. 26, No. 6,

pp. 1473-1480. ISSN: 0305-1048.

DOCUMENT TYPE:

Article English

LANGUAGE: ABSTRACT:

Modifications in the structure of a 260 bp DNA (hlyM) fragment from Escherichia coli caused by interaction with Pd(II) and Pt(II) complexes were studied. Cisplatin and transplatin (cis- and trans-PtCl2(NH3)2 respectively), Pt2Cl2(Spym)4 (SPYM = 2-mercaptopyrimidine anion), Pd-famotidine and Pt-famotidine were incubated with DNA for 24 h at 37degreeC and there observed with an atomic force microscope. Atomic

\*\*\*force\*\*\* microscopy (AFM) provides the opportunity for nanomater resolution in research on the interaction between nucleic acids and metal complexes. The complexes induced noticeable changes in DNA topography according to their different characteristics and structure. In the case of cisplatin a shortening in DNA strands was observed.

Transplatin and Pt2Cl2(SPYM)4 caused shortening and compaction, whilst an

aggregation of two strands was observed for the Pt-famotidine compound but not for the Pd-famotidine compound or the metal-free famotidine.

CONCEPT CODE: Biochemical Methods - Nucleic Acids, Purines and

CONCEPT CODE: Biochemical Methods Pyrimidines \*10052

Microscopy Techniques - General and Special Techniques

\*01052

Biochemical Methods - Minerals \*10059

Biophysics - Molecular Properties and Macromolecules

\*10506

INDEX TERMS: M

Major Concepts Biochemistry and Molecular Biophysics; Methods and

Techniques

INDEX TERMS: Chemicals & Biochemicals

cisplatin: Royston, quantitative analysis; thyM sets; palladium (II) ion: quantitative analysis; plutice of ion: quantitative analysis; transplatin: Royston, quantitative analysis; DNA: quantitative analysis

INDEX TERMS:

Methods & Equipment

atomic force microscopy:
microscopy method; polymerase chain reaction:

amplification method, sequencing techniques; GeneAmp PCR system 2400: Perkin-Elmer Cetus, equipment; Nanoscope III Multimode AFM: Digital Instrumentals Inc, equipment

Miscellaneous Descriptors

INDEX TERMS:

nucleic acid-metal interaction

```
16065-88-6 (FALNADIUM 111)
                    22542-10-5 (PLATINUM (II))
                    14913-33-8 (TRANSPLATIN)
    ANSWER 11 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS
                   1997:243950 BIOSIS
ACCESSION NUMBER:
                    PREV199799543153
DOCUMENT NUMBER:
                    DNA looping by Ku and the DNA-dependent
TITLE:
                    protein kinase.
                    Cary, Robert B.; Peterson, Scott R.; Wang, Jinting; Bear,
AUTHOR(S):
                    David G.; Bradbury, E. Morton; Chen, David J. (1)
                    (1) Life Sci. Div., Los Alamos Natl. Lab., Mail Stop M888,
CORPORATE SOURCE:
                    Los Alamos, NM 87545 USA
                 , -Proceedings of the National Academy of Sciences of the
SOURCE:
                    United States of America, (1997) Vol. 94, No. 9, pp.
                    4267-4272.
                    ISSN: 0027-8424.
                    Article
DOCUMENT TYPE:
                    English
LANGUAGE:
ABSTRACT:
The DNA-dependent protein kinase (DNA-PK) is required for
***DNA*** double-strand break (DSB) repair and immunoglobulingene
rearrangement and may play a role in the regulation of transcription. The
***DNA*** -PK holoenzyme is composed of three polypeptide subunits: the
          binding Ku70/86 heterodimer and an apprxeq 460-kDa catalytic
subunit (DNA-PKcs). DNA-PK has been hypothesized to
assemble at DNA DSBs and play structural as well as signal
transduction roles in DSB repair. Recent advances in atomic
             microscopy (AFM) have resulted in a technology capable
of producing high resolution images of native protein and proteinnucleic acid
complexes without staining or metal coating. The AFM provides a rapid and
direct means of probing the protein-nucleic acid interactions
responsible for DNA repair and genetic regulation. Here we have
employed AFM as well as electron microscopy to visualize Ku and
***DNA*** -PK in association with DNA. A significant number of
***DNA*** molecules formed loops in the presence of Ku. DNA looping
appeared to be sequence-independent and unaffected by the presence of
***DNA*** -PKcs. Gel filtration of Ku in the absence and the presence of
          indicates that Ku does not form nonspecific aggregates. We conclude
that, when bound to DNA, Ku is capable of self-association. These
findings suggest that Ku binding at DNA DSBs will result in Ku
self-association and a physical tethering of the proten DNA structure
CONCEPT CODE:
                    Biochemical Studies - Nucleic Acids, Purines and
                    Pyrimidines *10062
                    Biochemical Studies - Proteins, Peptides and Amino Acids
                    Biophysics - Molecular Properties and Macromolecules
                    *10506
                    Enzymes - Physiological Studies *10808
                    Major Concepts
INDEX TERMS:
                    Biochemistry and Molecular Biophysics; Enzymology
                    (Biochemistry and Molecular Biophysics)
                    Chemicals & Biochemicals
INDEX TERMS:
                    PROTEIN KINASE
INDEX TERMS:
                    Miscellaneous Descriptors
                    BIOCHEMISTRY AND BIOPHYSICS; DNA; DNA
                    -DEPENDENT PROTEIN KINASE; KU70/86 HETERODIMER; LOOPING
                    9026-43-1 (PROTEIN KINASE)
REGISTRY NUMBER:
     ANSWER 12 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS
ACCESSION NUMBER: 1996:150019 BIOSIS
                   PREV199698722154
DOCUMENT NUMBER:
                    Atomic force microscopy o:
TITLE:
```

15663-27-1 (CISPLATIN)

REGISTRY NUMBER:

long and short double-stranded, single-stranded and

triple-stranded nucleic acids.

Hansma, Helen G. (1); Revenko, Irene; Kim, Kery; Laney, AUTHOR(S):

Daniel E.

(1) Dep. Physics, Univ. California, Santa Barbara CA 93106 CORPORATE SOURCE:

USA

Nucleic Acids Research, (1996) Vol. 24, No. 4, pp. 713-720. SOURCE:

ISSN: 0305-1048.

DOCUMENT TYPE:

Article English

LANGUAGE: ABSTRACT:

Atomic force microscopy (AFM, also called

scanning force microscopy) is proving to be a useful

technique for imaging DNA. Thus it is important to push the limits of

AFM imaging in order to explore both what types of DNA can be

reliably imaged and identified and also what substrates and methods of sample

preparation are suitable. The following advances in AFM of DNA are

presented here. (i) DNA molecules as short as 25 bases can be seen by

AFM. The short single-stranded DNAs imaged here (25 and 50 bases long) appeared globular in the AFM, perhaps because they are all capable of intramolecular base pairing and because the DNAs were in a Mg(II) buffer, which facilitates intramolecular cross-bridging. (ii) AFM images in air of short double-stranded

molecules, 100-200 bp, gave lengths consistent with A-DNA

. (iii) AFM images of poly(A) show both short bent lumpy molecules with an apparent persistence length of 40 nm and long straight molecules with an apparent persistence length of 600 nm. For comparison, the apparent persistence

length for double-stranded DNA from vphi-X-174 under the same

conditions was 80 nm. (iv) Structures believed to be triple-stranded were seen in samples of poly(dA) cntdot poly(dT) and poly(dG)

cntdot poly(dC). These structures were twice as high as double-stranded

\*\*\*DNA\*\*\* and the same width. (v) Entire molecules of lambda DNA, apprx 16 mu-m long, were imaged clearly in overlapping stank. \*\*\*DNA\*\*\* was imaged on oxidized silicon, although less drearly than on miss.

Microscopy Techniques - General and Special Techniques CONCEPT CODE:

01052

Biochemical Methods - Nucleic Acids, Purines and

Pyrimidines \*10052

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biophysics - General Biophysical Techniques \*10504 Biophysics - Molecular Properties and Macromolecules

Physiology and Biochemistry of Bacteria \*31000

Genetics of Bacteria and Viruses \*31500

BIOSYSTEMATIC CODE: Bacteria - General Unspecified \*05000

INDEX TERMS:

Major Concepts

Biochemistry and Molecular Biophysics; Genetics; Methods

and Techniques; Physiology

INDEX TERMS:

Miscellaneous Descriptors

CROSS-BRIDGING; DNA IMAGING; INTRAMOLECULAR BASE

PAIRING; LAMBDA-DNA; PLASMID DNA;

SCANNING FORCE MICROSCOPY

ORGANISM:

Super Taxa

Bacteria - General Unspecified: Eupacteria, Bacteria

ORGANISM:

Organism Name

bacteria (Bacteria - General Unspecified)

ORGANISM: Organism Superterms

bacteria; eubacteria; microorganisms

ANSWER 13 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER:

1996:125528 BIOSIS

DOCUMENT NUMBER:

PREV199698697663

TITLE:

Scanning probe microscopy in microbiology.

AUTHOR(S):

Firtel, M.,; Beveridge, T. J. (1)

CORPORATE SOURCE:

(1) Department Microbiology, Fabrilty Medicine, Triversing

Toronto, Toronto, ON M5S 1A8 Canada

SOURCE:

Micron, (1995) Vol. 26, No. 4, pp. 347-362.

ISSN: 0968-4328.

DOCUMENT TYPE:

General Review

LANGUAGE:

English

CONCEPT CODE:

Microscopy Techniques - General and Special Techniques

Biochemical Methods - Nucleic Acids, Purines and

Pyrimidines \*10052

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines 10062

Biophysics - Molecular Properties and Macromolecules

\*10506

Anatomy and Histology, General and Comparative -

Microscopic and Ultramicroscopic Anatomy \*11108

Metabolism - Nucleic Acids, Purines and Pyrimidines \*13014

Morphology and Cytology of Bacteria \*30500 Physiology and Biochemistry of Bacteria

Genetics of Bacteria and Viruses \*31500

Microbiological Apparatus, Methods and Medula +30000

Microbiological Ultrastructure \* 32300

Virology - Bacteriophage \*33504

Virology - Animal Host Viruses \*33506 Virology - Plant Host Viruses \*33508

02600 BIOSYSTEMATIC CODE: Animal Viruses - General Bacterial Viruses - General

02800 Plant Viruses - General

Enterobacteriaceae 06702 07701 Deinococcaceae Methanomicrobiaceae 09531 Halobacteriaceae \*09711

INDEX TERMS:

INDEX TERMS:

Major Concepts

Biochemistry and Molecular Biophysics; Cell Biology;

Genetics; Metabolism; Methods and Techniques; Microbiology;

Morphology; Physiology Miscellaneous Descriptors

ANALYTICAL METHOD; ATOMIC FORCE

MICROSCOPY; BACTERIAL SURFACE LAYER; BACTERIOPHAGE

SURFACE LAYER; DNA; MICROBIAL CELL

VISCOELASTICITY; NUCLEIC ACID; SCANNING

TUNNELLING MICROSCOPY; SURFACE TOPOGRAPHY; VIHAL

SURFACE LAYER

ORGANISM:

Super Taxa

Animal Viruses - General: Viruses; Bacterial Viruses -General: Viruses; Deinococcaceae: Eubacteria, Bacteria; Enterobacteriaceae: Eubacteria, Bacteria; Halobacteriaceae:

Archaeobacteria, Bacteria; Methanomicrobiaceae:

Archaeobacteria, Bacteria; Plant Viruses - General: Viruses

ORGANISM:

Organism Name

animal viruses (Animal Viruses - General); bacterial viruses (Bacterial Viruses - General); Deinococcus

radiodurans (Deinococcaceae); Escherichia coli

(Enterobacteriaceae); Halobacterium (Halobacteriaceae); Methanospirillum hungatei (Methanomicrobiaceae); Plant

Viruses (Plant Viruses - General)

ORGANISM:

Organism Superterms

archaeobacteria; bacteria; eubacteria; microorganisms;

viruses

ANSWER 14 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

DOCUMENT NUMBER:

ACCESSION NUMBER: 1995:201818 BIOSIS PREV199598216118

TITLE:

A new DNA nanostructure, the G-wire, imaged by

scanning probe microscopy.

AUTHOR(S):

Marsh, Thomas C.; Vesenka, James; Henerson, Eric (1)

CORPORATE SOURCE:

(1) Dep. Zool. Genet., 2112 Mol. Biol. Build., Iowa State

Univ., Ames, IA 50011 USA

SOURCE:

Nucleic Acids Research, (1995) Vol. 23, No. 4, pp. 696-10-.

ISSN: 0305-1048.

DOCUMENT TYPE: LANGUAGE:

Article English

ABSTRACT:

G-DNA is a polymorphic family of quadruple helical nucleic acid structures containing quanine tetrad motifs (G-quartets; Williamsch, J.B., Raghuraman, M.K. and Cech, TR. (1989) Cell 59,871-880; Williamson, J.R. (1993, Proc. Natl. Acad Sci. USA 90, 3124-3124). Guanine rich oligonucleotides that are self-complimentary, as found in many telomeric G-strand repeat sequences,

form G-DNA In the presence of monovalent and/or divalent metal

cations. In this report we use the atomic force microscope (AFM) to explore the structural characteristics of long, linear polymers formed by the telomeric oligonucleotide d(GGGG TTGGGG) in the presence of specific metal cations. In the AFM these polymers, termed G-wires, appear as filaments whose height and length are determined by the metal ions present during the self-assembly process. The highly ordered, controllable self-assembly of G-wires could provide a basis for developing advanced biomaterials.

CONCEPT CODE:

Microscopy Techniques - Electron Microscopy (1918

Biochemical Methods - Nucleic Acids, Purines and

Pyrimidines 10052

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biophysics - General Biophysical Techniques Biophysics - Molecular Properties and Macromolecules

INDEX TERMS:

Mafor Concepts

Biochemistry and Molecular Biophysics; Methods and

Techniques

INDEX TERMS:

Miscellaneous Descriptors

ANALYTICAL METHOD; ATOMIC FORCE

MICROSCOPE; ION BIOSENSOR; OLIGONUCLEOTIDE

ANSWER 15 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

DOCUMENT NUMBER:

ACCESSION NUMBER: 1994:327879 BIOSIS PREV199497340879

TITLE:

Atomic force microscopy in

basic and applied nucleic acid research.

AUTHOR(S):

Henderson, Eric

CORPORATE SOURCE:

SOURCE:

Zool. and Genetics, Iowa State Univ., Ames, IA 50011 USA Clinical Chemistry, (1994) Vol. 40, No. 4, pp. 653. Meeting Info.: 8th San Diego Conference on Beyond DNA Probes San Diego, California, USA November 18-20, 1993

ISSN: 0009-9147.

DOCUMENT TYPE:

LANGUAGE:

English

Conference

CONCEPT CODE:

General Biology - Symposia, Transactions and Proceedings of

Conferences, Congresses, Review Annuals

Microscopy Techniques - General and Special Techniques

Genetics and Cytogenetics - General \*03502 Biochemical Methods - Nucleic Acids, Purines and

Pyrimidines \*10052

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biophysics - General Biophysical Techniques \*10504 Biophysics - Molecular Properties and Macromolecules

\*10506 BIOSYSTEMATIC CODE:

+00500

INDEX TERMS:

Major Concepts

Biochemistry and Molecular Biophysics; Genetics; Methods

and Techniques

INDEX TERMS:

Miscellaneous Descriptors

DNA TOPOGRAPHY; MEETING ABSTRACT; VISUALIZATION

METHOD

ORGANISM:

Organism Name

organisms (Organisms - Unspecified)

L3 ANSWER 16 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER:

1992:156807 BIOSIS

DOCUMENT NUMBER:

BR42:73007

TITLE:

ATOMIC FORCE MICROSCOPY

AUTHOR(S):

IMAGING OF LARGE DOUBLE STRANDED DNA MOLECULES.
LYUBCHENKO Y L; GALL A A; SHLYAKHTENKO L S; HARRINGTON R E;

LINDSAY S M

CORPORATE SOURCE:

DEP. BIOCHEMISTRY, UNIV. NEVADA RENO, RENO, NV 89557.

SOURCE:

JOINT MEETING OF THE AMERICAN SOCIETY FOR BIOCHEMISTRY AND MOLECULAR BIOLOGY/BIOPHYSICAL SOCIETY, HOUSTON, TEXAS, USA, FEBRUARY 9-13, 1992. FASEB (FED AM SOC EXP BIOL) J, 11992

6 (1), A149.

CODEN: FAJOEC. ISSN: 0892-6638.

DOCUMENT TYPE:

Conference

FILE SEGMENT: LANGUAGE:

BR; OLD English

CONCEPT CODE:

General Biology - Symposia, Transactions and Proceedings of

Conferences, Congresses, Review Annuals 30520 Genetics and Cytogenetics - General \*03502 Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biophysics - Molecular Properties and Macromolecules

\*10506

INDEX TERMS:

Miscellaneous Descriptors

ABSTRACT GENOMIC MAPPING NUCLEIC ACID-PROTEIN

COMPLEXES

ANSWER 1 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS L3

2000:544893 BIOSIS ACCESSION NUMBER: PREV200000544893 DOCUMENT NUMBER:

The effect of super-oxidized water on Escherichia coli. TITLE: Zinkevich, V. (1); Beech, I. B.; Tapper, R.; Bogdarina,

AUTHOR(S): (1) School of Pharmacy and Biomedical Sciences, University CORPORATE SOURCE: of Portsmouth, White Swan Road, St. Michael's Building,

Portsmouth, POI 2DT UK

Journal of Hospital Infection, (October, 2000) Vol. 46, We. SOURCE:

2, pp. 153-156. print.

ISSN: 0195-6701.

Article DOCUMENT TYPE: English LANGUAGE: English SUMMARY LANGUAGE:

ABSTRACT:

The mechanism of action of Sterilox, a non-toxic liquid biocide produced by electrolysis of a dilute saline solution, upon planktonic cells of Escherichia coli JM109 was investigated using protein and nucleic acid analysis. The results revealed total destruction of chromosomal and plasmid DNA

, RNA and proteins of E. coli within 5 min of exposure. Our earlier

investigation conducted using atomic force

\*\*\*microscopy\*\*\* imaging revealed swelling and rupture of E. coli cells with release of cytoplasm. We propose that the biocidal properties of Sterilox are due to its effect upon constituents of the bacterial cell including proteins and nucleic acids.

CONCEPT CODE:

Physiology and Biochemistry of Bacteria \*31000 Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines :10062

Biochemical Studies - Proteins, Peptides and Amino Acids

\*10064

Public Health: Environmental Health - Sewage Disposal and

Sanitary Measures \*37014

Pest Control, General; Pesticides; Herbicides \*54600

BIOSYSTEMATIC CODE: Enterobacteriaceae 06702

Major Concepts INDEX TERMS:

Sanitation

Chemicals & Biochemicals INDEX TERMS:

Sterilox: biocide; plasmid DNA; plasmid RNA;

proteins; super-oxidized water

Methods & Equipment INDEX TERMS:

electrolysis: synthetic method; nucleic acid

analysis: analytical method; protein analysis: analytical

method

Super Taxa ORGANISM:

Enterobacteriaceae: Facultatively Anaerobic Gram-Negative Rods, Eubacteria, Bacteria, Microorganisms; Hominidae: Primates, Mammalia, Vertebrata, Chordata, Animalia

Organism Name ORGANISM:

Escherichia coli (Enterobacteriaceae): pathogen,

strain-JM109; human (Hominidae): patient

Organism Superterms ORGANISM:

Animals; Bacteria; Chordates; Eubacteria; Humans; Mammals;

Microorganisms; Primates; Vertebrates

ANSWER 2 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS L3

2000:497584 BIOSIS ACCESSION NUMBER: PREV200000497705 DOCUMENT NUMBER:

Near field microscopies: From isolated molecules to living TITLE:

cells. Delain, E. (1); Michel, D. (1); Le Grimellec, Ch. AUTHOR(S):

(1) Laboratoire de Microscopie Moleculaire et CORPORATE SOURCE:

Cellulaire/CNRS UMR 8532, Institut Gustave Roussy, Rue

Camille Desmoulins, Villejuif France Morphologie, (Juin, 2000) Vol. 84, No. 265, pp. 25-30. SOURCE:

print.

ISSN: 1286-0115.

DOCUMENT TYPE:

Article French

LANGUAGE:

English; French

SUMMARY LANGUAGE: ABSTRACT:

Near field (or scanning probe) microscopy is a recent technology which, owing to the huge amount of publications, is becoming a reference method in molecular and cellular imaging. These microscopies consist in the scanning of the sample, line by line, with a very tiny tip and thus providing informations on its surface down to the nanometer scale. These methods gather scanning tunelling microscopy (STM), which measures a current between the tip and the specimen support, atomic force

\*\*\*microscopy\*\*\* (AFM), which measures the repulsive and attractive forces of the tip in contact or very close to the specimen, and scanning near field optical microscopies (SNOM), for which a glass tip allows to catch light signals. Atomic force microscopy, which allows

the observation of specimens in air or physiological conditions environments, is presently dominant in biology, in complementarity with the classical optimal and electron microscopies, which by the way, have also shown considerable improvements during the last years. The complementarity of these microscopies is due to their very different basic principles, which provide them various possibilities and limits. The biological applications of STM is limited by the need of conducting samples, but the different models of SNOM, often still is development, allow to consider very interesting applications, particularly for detecting very faint and tiny fluorescence signals. Different examples will be given concerning the visualization by AFM of isolated DNA molecules, naked or associated with proteins, the observation of intact or decondensed chromosomes, as well as living cells. One of the originality of AFM is its capacity to observed objects in a wide range of enlargements, with fields from

a few hundred of nanometers to several micrometers. CONCEPT CODE:

Cytology and Cytochemistry - General

Biochemical Studies - General \*10060

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biochemical Studies - Proteins, Peptides and Amino Acids

\*10064

INDEX TERMS:

Major Concepts

Biochemistry and Molecular Biophysics; Cell Biology;

Methods and Techniques

INDEX TERMS:

Parts, Structures, & Systems of Organisms

cells: near field microscopic study

INDEX TERMS:

Chemicals & Biochemicals

double stranded DNA: microscopic study; nucleic acids: microscopic study; proteins:

microscopic study

Methods & Equipment INDEX TERMS:

scanning near field optical microscopy: analytical method; scanning probe microscopy [near field microscopy]: analytical method

ORGANISM:

Super Taxa Organisms

ORGANISM:

Organism Name

organism (Organisms)

ANSWER 3 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: DOCUMENT NUMBER:

2000:309262 BIOSIS PREV200000309262

TITLE:

A dimer as a building block in assembling RNA: A hexamer

that gears bacterial virus phi29 DNA

-translocating machinery.

AUTHOR(S): SOURCE:

Chen, Chaoping; Sheng, Sitong; Shao, Zhifeng; Guo, Peixuan Journal of Biological Chemistry, (June 9, 2000) Vol. 275,

No. 23, pp. 17510-17516. print.

ISSN: 0021-9258.

DOCUMENT TYPE:

Article English

LANGUAGE: SUMMARY LANGUAGE:

English

ABSTRACT:

Six RNA (pRNA) molecules form a hexamer, via hand-in-hand interaction, to dear

bacterial virus phi29 DNA translocation machinery. Here we report the pathway and the conditions for the hexamer formation. Stable pRNA dimers and trimers were assembled in solution, isolated from native gels, and separated by sedimentation, providing a model system for the study of RNA dimers and trimers

in a protein-free environment. Cryoatomic force microscopy

revealed that monomers displayed a check-mark shaped outline, dimers exhibited an elongated shape, and trimers formed a triangle. Dimerization of pRNA was promoted by a variety of cations including spermidine, whereas procapsid

binding and DNA packaging required specific divalent cations,

including Mg2+, Ca2+, and Mn2+. Both the tandem and fused pRNA dimers with complementary loops designed to form even-numbered rings were active in

packaging, whereas those without complementary loops were inactive. \* \* \* DNA \* \* \* We conclude that dimers are the building blocks of the hexamer, and the pathway of building a hexamer is: dimer fwdarw tetramer fwdarw hexamer. The Hill coefficient of 2.5 suggests that there are three binding sites with cooperative

binding on the surface of the procapsid. The two interacting loops played a key role in recruiting the incoming dimer, whereas the procapsid served as the

foundation for hexamer assembly.

Biochemical Studies - Nucleic Acids, Purines and CONCEPT CODE:

Pyrimidines \*10062

Methods, Materials and Apparatus, General - Laboratory

Methods \*01004

Virology - Animal Host Viruses \*33506 Biochemical Studies - Minerals

INDEX TERMS:

Major Concepts

Biochemistry and Molecular Biophysics; Methods and

Techniques

INDEX TERMS:

Chemicals & Biochemicals

DNA-translocating machinery; RNA: assay,

isolation, separation, synthesis; calcium (II); magnesium

(II); manganese (II)

INDEX TERMS:

Methods & Equipment PAGE [polyacrylamide gel electrophoresis]: gel

electrophoresis, isolation method; RNA binding assay: analytical method, binding assays; RNA synthesis: nucleic acid synthesis, synthetic method; cryo-

atomic force microscopy

[cryo-AFM]: microscopy method, microscopy : CB; dimer binding competition assay:

Analysis/Characterization Techniques: CB, analytical method; sucrose gradient sedimentation: Extraction, Isolation, Purification and Separation Techniques,

separation method; virion assembly assay:

Analysis/Characterization Techniques: CB, analytical method

Super Taxa ORGANISM:

Podoviridae: Bacterial Viruses, Viruses, Microorganisms

Organism Name ORGANISM:

> phi29 (Podoviridae) Organism Superterms

Bacterial Viruses; Microorganisms; Viruses

REGISTRY NUMBER:

14127-61-8 (CALCIUM (II)) 22537-22-0 (MAGNESIUM (II); 16397-91-4 (MANGANESE (II))

ANSWER 4 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS ACCESSION NUMBER: DOCUMENT NUMBER:

2000:246496 BIOSIS PREV200000246496

TITLE:

L3

ORGANISM:

PNA-dependent gene chemistry: Stable coupling or peptides

and oligonucleotides to plasmid DNA.

Zelphati, O.; Liang, X.; Nguyen, C.; Barlow, S.; Sheng, S.; AUTHOR(S):

Shao, Z.; Felgner, P. L. (1)

(1) Gene Therapy Systems, 10190 Telesis Court, San Diego, CORPORATE SOURCE:

CA, 92121 USA

Biotechniques, (Feb., 2000) Vol. 28, No. 2, pp. 304-316. SOURCE:

ISSN: 0736-6205.

DOCUMENT TYPE:

Article Enalish English

SUMMARY LANGUAGE:

ABSTRACT:

LANGUAGE:

Two approaches are described for stably conjugating peptides, proteins and oligonucleotides onto plasmid DNA. Both methods use a peptide

\*\*\*nucleic\*\*\* acid (PNA) clamp, which binds irreversibly and specifically to

a binding site cloned into the plasmid. The first approach uses a

biotin-conjugated PNA clamp that can be used to introduce functional biotin

groups onto the plasmid to which streptavidin can bind. Atomic microscopy images of linearized plasmid show

\*\*\*force\*\*\* streptavidin localized at the predicted PNA binding site on the DNA strand. Peptides and oligonucleotides containing free thiol groups were conjugated to maleimide streptavidin, and these streptavidin conjugates were bound to the biotin-PNA-labeled plasmid. In this way, peptides and

oligonucleotides could be brought into stable association with the plasmid. A second approach used a maleimide-conjugated PNA clamp. Methods are described for conjugating thiclated peptides and oligonucleotides directly to the maleimide-PNA-DNA hybrid. This straightforward technology offers an

easy approach to introduce functional groups onto plasmid DNA without disturbing its transcriptional activity.

Biochemical Methods - Nucleic Acids, Purines and CONCEPT CODE:

Pyrimidines \*10052

Biochemical Studies - Proteins, Peptides and Amino Acids

Biophysics - General Biophysical Techniques \*10504

Genetics of Bacteria and Viruses \*31500

INDEX TERMS: Major Concepts

Molecular Genetics (Biochemistry and Molecular Biophysics;

Methods and Techniques

Chemicals & Biochemicals INDEX TERMS:

oligonucleotides; peptide nucleic acid; peptides;

plasmid DNA

INDEX TERMS: Methods & Equipment

ATTO-TAG labeling kit: Molecular Probes, equipment; agarcse

gel electrophoresis: analytical method, gel

electrophoresis; atomic force microscopy: microscopy method, microscopy: CB, microscopy: CT;

transfection: gene expression/vector techniques, genetic

method; transmission electron microscopy:

electron microscopy: CB, electron microscopy: CT, microscopy method

INDEX TERMS:

Miscellaneous Descriptors PNA-dependent gene chemistry

ANSWER 5 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS L3

ACCESSION NUMBER: DOCUMENT NUMBER:

2000:61886 BIOSIS PREV200000061886

TITLE:

Polymerase activities and RNA structures in the

atomic force microscope.

AUTHOR(S):

Hansma, Helen G. (1); Golan, Roxana; Hsieh, Wan;

CORPORATE SOURCE:

Daubendiek, Sarah L.; Kool, Eric T. (1) Department of Physics, University of California, Santa

Barbara, Santa Barbara, CA USA

SOURCE:

Journal of Structural Biology, (Oct., 1999) Vol. 127, No.

3, pp. 240-247.

ISSN: 1047-8477.

DOCUMENT TYPE:

Article English

LANGUAGE: SUMMARY LANGUAGE:

English

ABSTRACT:

The structures of the reaction products are the basis for novel polymerase

assays using the atomic force microscope (AFM). Polymerases

are the enzymes involved in transcription and replication of DNA. Rapid semiquantitative estimates of the activity of DNA polymerases

such as Sequenase, Taq polymerase, and AMV reverse transcriptase and RNA

polymerases (RNAP) such as Escherichia coli RNAP were obtained from AFM images

of the nucleic acids after polymerase reactions. DNA

polymerases were assayed via replication of the single-stranded PHIX-174 virion. RNAP was assayed via transcription, using a rolling circle DNA

template that produces long strands of RNA. In some cases, AFM was better than

agarose gel electrophoresis for assaying DNA polymerase activity, since aggregation prevented the DNA from entering the agarose gel.

Extended molecules of single-stranded RNA synthesized with the rolling circle template showed varied conformations and degrees of stretching. \* \* \* DNA \* \* \*

Some structural differences were observed between two RNAs-a ribozyme

concatamer and an RNA with 90% purines.

Genetics and Cytogenetics - General \*03502 CONCEPT CODE:

Microscopy Techniques - Electron Microscopy \*01058

Biochemical Methods - Nucleic Acids, Purines and

Pyrimidines **\***10052 Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Replication, Transcription, Translation \*10300 Biophysics - Molecular Properties and Macromolecules

\*10506

Enzymes - Methods \*10804

Enzymes - Chemical and Physical \*10806

Physiology and Biochemistry of Bacteria \*31000

Genetics of Bacteria and Viruses \*31500 Virology - General; Methods \*33502

BIOSYSTEMATIC CODE: Microviridae 02706

Enterobacteriaceae 06702

INDEX TERMS:

Major Concepts

Enzymology (Biochemistry and Molecular Biophysics); Molecular Genetics (Biochemistry and Molecular Biophysics

INDEX TERMS:

Chemicals & Biochemicals

AMV reverse transcriptase; DNA: rolling circle template; DNA polymerase; RNA: structure; RNA

polymerase; Taq polymerase

INDEX TERMS:

Methods & Equipment

atomic force microscope: laboratory

equipment; atomic force

microscopy: analytical method, microscopy

: CB, molecular imaging method

ORGANISM:

Super Taxa

Enterobacteriaceae: Facultatively Anaerobic Gram-Negative Rods, Eubacteria, Bacteria, Microorganisms; Microviridae:

Bacterial Viruses, Viruses, Microorganisms

ORGANISM:

Organism Name

Escherichia coli (Enterobacteriaceae); bacteriophage

phi-X-174 (Microviridae)

ORGANISM:

Organism Superterms

Bacteria; Bacterial Viruses; Eubacteria; Microorganisms;

Viruses

REGISTRY NUMBER:

9012-90-2 (**DNA** POLYMERASE)

9014-24-8 (RNA POLYMERASE)

ANSWER 6 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER:

2000:2359 BIOSIS

DOCUMENT NUMBER:

PREV200000002359

TITLE:

DNA toroids: Stages in condensation.

AUTHOR(S):

Golan, Roxana; Pietrasanta, Lia I.; Hsieh, Wan; Hansma,

Helen G. (1)

CORPORATE SOURCE:

(1) Department of Physics, UCSB, Santa Barbara, CA, 93106

SOURCE:

Biochemistry, (Oct. 19, 1999) Vol. 38, No. 42, pp.

14069-14076.

ISSN: 0006-2960.

DOCUMENT TYPE:

Article

LANGUAGE:

English

SUMMARY LANGUAGE:

English

ABSTRACT:

The effects of polylysine (PLL) and PLL-asialoorosomucoid (AsOR) on DNA

condensation have been analyzed by AFM. Different types of condensed structures were observed, which show a sequence of conformational

changes as circular plasmid DNA molecules condense progressively. The structures range from circular molecules with the length of the plasmid

to small toroids and short rods with apprx1/6 to 1/8 the contour \* \* \* DNA \* \* \* length of the uncondensed circular DNA. Single plasmid molecules of

6800 base pairs (bp) condense into single toroids of apprx110 nm diameter,

measured center-to-center. The results are consistent with a mode! for condensation in which circular DNA molecules fold several \* \* \* DNA \* \* \*

times into progressively shorter rods. Structures intermediate between toroids and rods suggest that at least some toroids may form by the opening up of rods

as proposed by Dunlap et al. ((1997) Nucleic Acids Res. 25, 3095).

Toroids and rods formed at lysine: nucleotide ratios of 5:1 and 6:1. This high lys ine: nucleotide ratio is discussed in relation to entropic considerations and the overcharging of macroions. PLL-AsOR is much more effective than PLL

alone for condensing DNA, because several PLL molecules are attached to a single AsOR molecule, resulting in an increased cation density.

CONCEPT CODE:

Genetics and Cytogenetics - General \*03502

Microscopy Techniques - General and Special Techniques

Biochemical Methods - General \*10050 Biochemical Studies - General \*10060

Biophysics - General Biophysical Studies \*10502

INDEX TERMS:

INDEX TERMS:

Major Concepts

Molecular Genetics (Biochemistry and Molecular Biophysics);

Methods and Techniques Chemicals & Biochemicals

DNA toroids: analysis, condensation stages;

polylysine: DNA condenser; polylysine-

asialoorosomucoid: DNA condenser

INDEX TERMS:

Methods & Equipment

Fast Flow Q Sepharose anion exchange chromatography:

chromatographic techniques, separation method;

atomic force microscopy:

microscopy method, microscopy: CB

REGISTRY NUMBER:

25104-18-1Q (POLYLYSINE) 38000-06-5Q (POLYLYSINE)

ACCESSION NUMBER:

ANSWER 7 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS 1999:187181 BIOSIS

DOCUMENT NUMBER:

PREV199900187181

TITLE:

Activity of a single exonuclease revealed by atomic

force microscopy.

AUTHOR(S):

Takeuchi, M. (1); Okada, T. (1)

CORPORATE SOURCE:

(1) Joint Research Center for Atom Technology c/o NAIR,

1-1-4 Higashi, Tsukuba, Ibaraki, 305-0046 Japan

SOURCE:

Biophysical Journal, (Jan., 1999) Vol. 76, No. 1 PART 2,

pp. A132.

Meeting Info.: Forty-third Annual Meeting of the Biophysical Society Baltimore, Maryland, USA February 13-17, 1999

ISSN: 0006-3495.

DOCUMENT TYPE:

Conference

LANGUAGE:

English

CONCEPT CODE:

\*10060 Biochemical Studies - General Biochemical Methods - General \*10050

Enzymes - General and Comparative Studies; Coenzymes

General Biology - Symposia, Transactions and Proceedings of

Conferences, Congresses, Review Annuals \*00520

INDEX TERMS:

Major Concepts

Enzymology (Biochemistry and Molecular Biophysics); Methods

and Techniques

INDEX TERMS:

Chemicals & Biochemicals exonuclease: activity; DNA

INDEX TERMS:

Methods & Equipment

atomic force microscopy:

microscopy method, microscopy: CB

INDEX TERMS:

Miscellaneous Descriptors

enzyme kinetics; protein-nucleic acid

interactions; Meeting Abstract; Meeting Poster

REGISTRY NUMBER:

37228-74-3 (EXONUCLEASE)

ANSWER 8 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: DOCUMENT NUMBER:

1998:256680 BIOSIS PREV199800256680

TITLE:

Novel vectors for gene delivery formed by self-assembly of

DNA with poly(L-lysine) grafted with hydrophilic

polymers.

AUTHOR(S):

Toncheva, Veska; Wolfert, Margreet A.; Dash, Philip R.; Oupicky, David; Ulbrich, Karel; Seymour, Leonard W. (1);

Schacht, Etienne H.

CORPORATE SOURCE:

(1) CRC Inst. Cancer Studies, Univ. Birmingham, Birmingham

B15 2TA UK

SOURCE:

Biochimica et Biophysica Acta, (May 8, 1998) Vol. 138, No.

3, pp. 354-368. ISSN: 0006-3002.

DOCUMENT TYPE:

Article

LANGUAGE:

English

ABSTRACT:

Complexes formed between DNA and cationic polymers are attracting increasing attention as novel synthetic vectors for delivery of genes. We are trying to improve biological properties of such complexes by oriented self-assembly of DNA with cationic-hydrophilic block copolymers, designed to enshroud the complex within a protective hydrophilic polymer corona. Poly(L-lysine) (pLL) grafted with range of hydrophilic polymer blocks, including poly(ethylene glycol) (pEG), dextran and poly(N-(2hydroxypropyl)methacrylamide) (pHPMA), shows efficient binding to DNA and mediates particle self-assembly and inhibition of ethidium bromide/ fluorescence. The complexes formed are discrete and typically about 100 nm diameter, viewed by atomic force microscopy

. Surface charges are slightly shielded by the presence of the hydrophilic polymer, and complexes generally show decreased cytotoxicity compared with simple pLL/DNA complexes. pEG-containing complexes show increased transfection activity against cells in vitro. Complexes formed with all polymer conjugates showed greater aqueous solubility than simple pLL/DNA complexes, particularly at charge neutrality. These materials appear to have the ability to regulate the physicochemical and biological properties of polycation/DNA complexes, and should find important applications in packaging of nucleic acids for specific biological applications. Genetics and Cytogenetics - General \*03502 CONCEPT CODE:

Microscopy Techniques - Electron Microscopy \*C1058 Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biochemical Studies - General \*10060

Major Concepts INDEX TERMS:

Biochemistry and Molecular Biophysics

INDEX TERMS:

Chemicals & Biochemicals

hydrophilic polymers; poly(L-lysine); DNA:

self-assembly

INDEX TERMS:

Methods & Equipment

atomic force microscopy: analytical method

INDEX TERMS:

Miscellaneous Descriptors gene delivery: novel vectors

REGISTRY NUMBER:

25104-18-1Q (POLY(L-LYSINE)) 38000-06-5Q (POLY(L-LYSINE))

ANSWER 9 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: DOCUMENT NUMBER:

1998:256659 BIOSIS PREV199800256659

TITLE:

Analysis of various sequence-specific triplexes by electron

and atomic force microscopies.

AUTHOR(S):

Cherny, Dimitry I. (1); Fourcade, Alain; Svinarchuk, Fedro;

Nielsen, Peter E.; Malvy, Claude; Delain, Etienne

CORPORATE SOURCE:

(1) Lab. Microscopie Cellularie Molecularie, URA 147, CNRS,

Inst. Gustave-Roussy, rue Camillie Desmoulin, F-94805

Villejuif France

SOURCE:

Biophysical Journal, (Feb., 1998) Vol. 74, No. 2 PART 1,

pp. 1015-1023. ISSN: 0006-3495.

DOCUMENT TYPE:

Article

LANGUAGE:

English

ABSTRACT:

Sequence-specific interactions of 20-mer G, A-containing triple herix-forming oligonucleotides (TFOs) and bis-PNAs (peptide nucleic acids) with

double-stranded DNA was visualized by electron (EM) and

force (AFM) microscopies. Triplexes formed by \*\*\*atomic\*\*\*

biotinylated TFOs are easily detected by both EM and AFM in which streptavidin is a marker. AFM images of the unlabeled triplex within a long plasmid

show a apprx0.4-nm height increment of the double helix within the \*\*\*DNA\*\*\* target site position. TFOs conjugated to a 74-nt-long oligonucleotide forming a 33-bp-long hairpin form extremely stable triplexes with the target site that are readily imaged by both EM and AFM as protruding DNA. The short duplex protrudes in a perpendicular direction relative to the double nelix axis, either in the plane of the support or out of it. In the latter case, the apparent height of the protrusion is apprx 1.5 nm, when that of the triplex site is increased by 0.3-0.4 nm. Triplex formation by bis-PNA, in which two decamers of PNA are connected via a flexible linker, causes deformations of the double helix at the target site, which is readily detected as kinks by both FM

and AFM. Moreover, AFM shows that these kinks are often accompanied by an increase in the DNA apparent height of apprx 35%. This work shows the first direct visualization of sequence-specific interaction of TFOs and PNAs, with their target sequences within long plasmid DNAs, through the measurements

of the apparent height of the DNA double helix by AFM.

\*03502 Genetics and Cytogenetics - General CONCEPT CODE:

Microscopy Techniques - Electron Microscopy \*01058 Biochemical Methods - Nucleic Acids, Purines and

Pyrimidines \*10052

Biochemical Methods - Proteins, Peptides and Amino Acids

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biochemical Studies - Proteins, Peptides and Amino Acids

Biophysics - Molecular Properties and Macromolecules

\*10506

INDEX TERMS:

Major Concepts

Methods and Techniques; Molecular Genetics (Biochemistry

and Molecular Biophysics) Chemicals & Biochemicals

bis-peptide nucleic acid: DNA

interaction; double-stranded DNA; triple

helix-forming oligonucleotide: DNA interaction

Methods & Equipment INDEX TERMS:

atomic force microscopy:

analytical method; electron microscopy:

analytical method

ANSWER 10 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS L3

ACCESSION NUMBER: DOCUMENT NUMBER:

1998:221880 BIOSIS PREV199800221880

TITLE:

Study of the interaction of DNA with displatin and other Pd(II) and Pt(II) complexes by atomic

force microscopy.

AUTHOR(S):

INDEX TERMS:

Onoa, G. Bibiana; Cervantes, Gemma; Moreno, Virtudes (1);

Prieto, M. Jose

CORPORATE SOURCE:

(1) Dep. Quim. Inorg., Gniv. Barcelona, Diagonal 647,

08028-Barcelona Spain

SOURCE:

Nucleic Acids Research, (March 15, 1998) Vol. 26, No. 6,

pp. 1473-1480. ISSN: 0305-1048.

DOCUMENT TYPE: LANGUAGE:

Article English

ABSTRACT:

Modifications in the structure of a 260 bp DNA (hlyM) fragment from Escherichia coli caused by interaction with Pd(II) and Pt(II) complexes were studied. Cisplatin and transplatin (cis- and trans-PtCl2(NH3)2 respectively;, Pt2Cl2(Spym)4 (SPYM = 2-mercaptopyrimidine anion), Pd-famotidine and Pt-famotidine were incubated with DNA for 24 h at 37degreeC and then

observed with an atomic force microscope. Atomic \*\*\*force\*\*\* microscopy (AFM) provides the opportunity for nanometer resolution in research on the interaction between nucleic acids and metal complexes. The complexes induced noticeable changes in DNA

topography according to their different characteristics and structure. In the case of displatin a shortening in DNA strands was observed.

Transplatin and Pt2Cl2(SPYM)4 caused shortening and compaction, whilst an aggregation of two strands was observed for the Pt-famotidine compound but not for the Pd-famotidine compound or the metal-free famotidine.

CONCEPT CODE:

Biochemical Methods - Nucleic Acids, Purines and

Pyrimidines \*10052

Microscopy Techniques - General and Special Techniques

\*01052

Biochemical Methods - Minerals \*10059

Biophysics - Molecular Properties and Macromolecules

\*10506

INDEX TERMS:

Major Concepts

Biochemistry and Molecular Biophysics; Methods and

Techniques

INDEX TERMS:

Chemicals & Biochemicals

cisplatin: Royston, quantitative analysis; hlyM gene; palladium (II) ion: quantitative analysis; platinum [11] ion: quantitative analysis; transplatin: Royston, quantitative analysis; DNA: quantitative analysis

Methods & Equipment INDEX TERMS:

atomic force microscopy:

microscopy method; polymerase chain reaction:

amplification method, sequencing techniques; GeneAmp PCR system 2400: Perkin-Elmer Cetus, equipment; Nanoscope III

Multimode AFM: Digital Instrumentals Inc, equipment Miscellaneous Descriptors

INDEX TERMS:

nucleic acid-metal interaction

REGISTRY NUMBER:

15663-27-1 (CISPLATIN)

16065-88-6 (PALLADIUM (II)) 22542-10-5 (PLATINUM (II)) 14913-33-8 (TRANSPLATIN)

L3 ANSWER 11 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: DOCUMENT NUMBER:

1997:243950 BIOSIS PREV199799543153

TITLE:

DNA looping by Ku and the DNA-dependent

protein kinase.

AUTHOR(S):

Cary, Robert B.; Peterson, Scott R.; Wang, Jinting; Bear,

David G.; Bradbury, E. Morton; Chen, David J. (1)

CORPORATE SOURCE:

(1) Life Sci. Div., Los Alamos Natl. Lab., Mail Stop M894,

Los Alamos, NM 87545 USA

SOURCE:

Proceedings of the National Academy of Sciences of the United States of America, (1997) Vol. 94, No. 9, pp.

4267-4272.

ISSN: 0027-8424.

DOCUMENT TYPE:

Article English

LANGUAGE: ABSTRACT:

The DNA-dependent protein kinase (DNA-PK) is required for

\*\*\*DNA\*\*\* double-strand break (DSB) repair and immunoglobulin gene rearrangement and may play a role in the regulation of transcription. The \*\*\*DNA\*\*\* -PK holoenzyme is composed of three polypoptide subunits: the

\*\*\*DNA\*\*\* binding Ku70/86 heterodimer and an apprxeq 460-kDa catalytic

subunit (DNA-PKcs). DNA-PK has been hypothesized to

assemble at DNA DSBs and play structural as well as signal

transduction roles in DSB repair. Recent advances in **atomic**\*\*\*force\*\*\* **microscopy** (AFM) have resulted in a technology capable

of producing high resolution images of native protein and proteinnucleic acid complexes without staining or metal coating. The AFM provides a rapid and

direct means of probing the protein-nucleic acid interactions responsible for **DNA** repair and genetic regulation. Here we have employed AFM as well as electron microscopy to visualize Ku and

\*\*\*DNA\*\*\* -PK in association with **DNA**. A significant number of \*\*\*DNA\*\*\* molecules formed loops in the presence of Ku. **DNA** looping

appeared to be sequence-independent and unaffected by the presence of

\*\*\*DNA\*\*\* -PKcs. Gel filtration of Ku in the absence and the presence of \*\*\*DNA\*\*\* indicates that Ku does not form nonspecific aggregates. We conclude

that, when bound to **DNA**, Ku is capable of self-association. These findings suggest that Ku binding at **DNA** DSBs will result in Ku self-association and a physical tethering of the broken **DNA** strands.

CONCEPT CODE:

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biochemical Studies - Proteins, Feptides and Amino Acids

\*10064

Biophysics - Molecular Properties and Macromolecules

\*10506

Enzymes - Physiological Studies \*10808

INDEX TERMS:

Major Concepts

Biochemistry and Molecular Biophysics; Enzymology

(Biochemistry and Molecular Biophysics)

INDEX TERMS:

Chemicals & Biochemicals

PROTEIN KINASE

INDEX TERMS:

Miscellaneous Descriptors

BIOCHEMISTRY AND BIOPHYSICS; DNA; DNA

-DEPENDENT PROTEIN KINASE; KU70/86 HETERODIMER; LOOPING

REGISTRY NUMBER: 9026-43-1 (PROTEIN KINASE)

L3 ANSWER 12 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: DOCUMENT NUMBER:

1996:150019 BIOSIS PREV199698722154

TITLE:

Atomic force microscopy of

long and short double-stranded, single-stranded and

triple-stranded nucleic acids.

Hansma, Helen G. (1); Revenko, Irene; Kim, Kery; Laney, AUTHOR(S):

Daniel E.

(1) Dep. Physics, Univ. California, Santa Barbara CA 93116 CORPORATE SOURCE:

Nucleic Acids Research, (1996) Vol. 24, No. 4, pp. 713-723. SOURCE:

ISSN: 0305-1048.

DOCUMENT TYPE:

Article English

LANGUAGE: ABSTRACT:

Atomic force microscopy (AFM, also called

scanning force microscopy) is proving to be a useful technique for imaging DNA. Thus it is important to push the limits of

AFM imaging in order to explore both what types of DNA can be

reliably imaged and identified and also what substrates and methods of sample

preparation are suitable. The following advances in AFM of DNA are

presented here. (i) DNA molecules as short as 25 bases can be seen by

AFM. The short single-stranded DNAs imaged here (25 and 50 bases long) appeared globular in the AFM, perhaps because they are all capable of intramolecular base pairing and because the DNAs were in a Mg(II) buffer, which facilitates

intramolecular cross-bridging. (ii) AFM images in air of short double-stranded

molecules, 100-200 bp, gave lengths consistent with A-DNA \*\*\*DNA\*\*\*

. (iii) AFM images of poly(A) show both short bent lumpy molecules with an apparent persistence length of 40 nm and long straight molecules with an apparent persistence length of 600 nm. For comparison, the apparent persistence

length for double-stranded DNA from vphi-X-174 under the same

conditions was 80 nm. (iv) Structures believed to be triple-stranded

\*\*\*DNA\*\*\* were seen in samples of poly(dA) cntdot poly(dT) and poly(dG)

entdot poly(dC). These structures were twice as high as double-stranded

\*\*\*DNA\*\*\* and the same width. (v) Entire molecules of lambda DNA, apprx 16 mu-m long, were imaged clearly in overlapping scans. (vi) Plasmid

\*\*\*DNA\*\*\* was imaged on oxidized silicon, although less clearly than on mica.

Microscopy Techniques - General and Special Techniques CONCEPT CODE:

01052

Biochemical Methods - Nucleic Acids, Purines and

Pyrimidines \*10052

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biophysics - General Biophysical Techniques \*10504 Biophysics - Molecular Properties and Macromolecules

\*10506

Physiology and Biochemistry of Bacteria

Genetics of Bacteria and Viruses \*31500

BIOSYSTEMATIC CODE: Bacteria - General Unspecified

INDEX TERMS:

Major Concepts

Biochemistry and Molecular Biophysics; Genetics; Methods

and Techniques; Physiology

INDEX TERMS:

Miscellaneous Descriptors

CROSS-BRIDGING; DNA IMAGING; INTRAMOLECULAR BASE

PAIRING; LAMBDA-DNA; PLASMID DNA;

SCANNING FORCE MICROSCOPY

ORGANISM:

Super Taxa

Bacteria - General Unspecified: Eubacteria, Bacteria

ORGANISM:

Organism Name bacteria (Bacteria - General Unspecified)

Organism Superterms ORGANISM:

bacteria; eubacteria; microorganisms

ANSWER 13 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER:

1996:125528 BIOSIS

DOCUMENT NUMBER:

PREV199698697663

TITLE:

Scanning probe microscopy in microbiology.

AUTHOR(S):

Firtel, M.,; Beveridge, T. J. (1)

CORPORATE SOURCE:

(1) Department Microbiology, Faculty Medicine, University

Toronto, Toronto, ON M5S 1A8 Canada

SOURCE:

Micron, (1995) Vol. 26, No. 4, pp. 347-362.

ISSN: 0968-4328.

DOCUMENT TYPE:

General Review English

LANGUAGE: CONCEPT CODE:

Microscopy Techniques - General and Special Techniques

\*01052

Biochemical Methods - Nucleic Acids, Purines and

\*10052 Pyrimidines

Biochemical Studies - Nucleic Acids, Purines and

10062 Pyrimidines

Biophysics - Molecular Properties and Macromolecules

Anatomy and Histology, General and Comparative -Microscopic and Ultramicroscopic Anatomy \*11108

Metabolism - Nucleic Acids, Purines and Pyrimidines

Morphology and Cytology of Bacteria \*30500 Physiology and Biochemistry of Bacteria

Genetics of Bacteria and Viruses \*31500

Microbiological Apparatus, Methods and Media \*32000

\*32300 Microbiological Ultrastructure

Virology - Bacteriophage \*33504

Virology - Animal Host Viruses \*33506 Virology - Plant Host Viruses \*33508

BIOSYSTEMATIC CODE: Animal Viruses - General 02600 02700 Bacterial Viruses - General 02800 Plant Viruses - General

> Enterobacteriaceae 96702 Deinococcaceae 07701 09531 Methanomicrobiaceae \*09711 Halobacteriaceae

INDEX TERMS:

INDEX TERMS:

Major Concepts

Biochemistry and Molecular Biophysics; Cell Biology;

Genetics; Metabolism; Methods and Techniques; Microbiology;

Morphology; Physiology

Miscellaneous Descriptors ANALYTICAL METHOD; ATOMIC FORCE

MICROSCOPY; BACTERIAL SURFACE LAYER; BACTERIOPHAGE

SURFACE LAYER; DNA; MICROBIAL CELL VISCOELASTICITY; NUCLEIC ACID; SCANNING

TUNNELLING MICROSCOPY; SURFACE TOPOGRAPHY; VIRAL

SURFACE LAYER

ORGANISM:

Super Taxa Animal Viruses - General: Viruses; Bacterial Viruses -General: Viruses; Deinococcaceae: Eubacteria, Bacteria; Enterobacteriaceae: Eubacteria, Bacteria; Halobacteriaceae:

Archaeobacteria, Bacteria; Methanomicrobiaceae:

Archaeobacteria, Bacteria; Plant Viruses - General: Viruses

ORGANISM:

Organism Name

animal viruses (Animal Viruses - General); bacterial viruses (Bacterial Viruses - General); Deinococcus

radiodurans (Deinococcaceae); Escherichia coli

(Enterobacteriaceae); Halobacterium (Halobacteriaceae); Methanospirillum hungatei (Methanomicrobiaceae,; Plant

Viruses (Plant Viruses - General)

ORGANISM:

Organism Superterms archaeobacteria; bacteria; eubacteria; microorganisms;

viruses

ANSWER 14 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: DOCUMENT NUMBER:

1995:201818 BIOSIS PREV199598216118

TITLE:

A new DNA nanostructure, the G-wire, imaged by

scanning probe microscopy.

AUTHOR(S):

Marsh, Thomas C.; Vesenka, James; Henerson, Eric (1)

CORPORATE SOURCE:

(1) Dep. Zool. Genet., 2112 Mol. Biol. Build., Iowa State

Univ., Ames, IA 50011 USA

SOURCE:

Nucleic Acids Research, (1995) Vol. 23, No. 4, pt. 696-10.

ISSN: 0305-1048.

DOCUMENT TYPE:

Article English

LANGUAGE: ABSTRACT:

G-DNA is a polymorphic family of quadruple helical nucleic acid structures containing guanine tetrad motifs (G-quartets; Williamson, J.R. Raghuraman, M.K. and Cech, TR. (1989) Cell 59,871-880; Williamson, J.R. (1993; Proc. Natl. Acad Sci. USA 90, 3124-3124). Guanine rich oligonucleotides that are self-complimentary, as found in many telomeric G-strand repeat sequences, form G. DNA In the presence of monovalent and/or divalent metal

form G-DNA In the presence of monovalent and/or divalent metal cations. In this report we use the **atomic force** microscope

(AFM) to explore the structural characteristics of long, linear polymers formed by the telomeric oligonucleotide d(GGGG TTGGGG) in the presence of specific metal cations. In the AFM these polymers, termed G-wires, appear as filaments whose height and length are determined by the metal ions present during the self-assembly process. The highly ordered, controllable self-assembly of G-wires could provide a basis for developing advanced biomaterials.

CONCEPT CODE:

Microscopy Techniques - Electron Microscopy \*01058

Biochemical Methods - Nucleic Acids, Purines and

Pyrimidines 10052

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biophysics + General Biophysical Techniques 10504 Biophysics - Molecular Properties and Macromolecules

\*10506

INDEX TERMS:

Major Concepts

Biochemistry and Molecular Biophysics; Methods and

Techniques

INDEX TERMS:

Miscellaneous Descriptors

ANALYTICAL METHOD; ATOMIC FORCE

MICROSCOPE; ION BIOSENSOR; OLIGONUCLEOTIDE

L3 ANSWER 15 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER: DOCUMENT NUMBER:

1994:327879 BIOSIS PREV199497340879

TITLE:

Atomic force microscopy in

basic and applied nucleic acid research.

AUTHOR(S):

Henderson, Eric

CORPORATE SOURCE:

SOURCE:

Zool. and Genetics, Iowa State Univ., Ames, IA 50011 USA Clinical Chemistry, (1994) Vol. 40, No. 4, pp. 653.
Meeting Info.: 8th San Diego Conference on Beyond DNA
Deather Car Diego Colifornia, USA November 18-20, 1993

Probes San Biego, California, USA November 18-20, 1993

ISSN: 0009-9147.

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CONCEPT CODE:

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\*01052

Genetics and Cytogenetics - General \*03502 Biochemical Methods - Nucleic Acids, Purines and

Pyrimidines \*10052

Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biophysics - General Biophysical Techniques \*10504 Biophysics - Molecular Properties and Macromolecules

\*10506

BIOSYSTEMATIC CODE:

\*00500

INDEX TERMS:

Major Concepts

Biochemistry and Molecular Biophysics; Genetics; Methods

and Techniques

INDEX TERMS:

Miscellaneous Descriptors

DNA TOPOGRAPHY; MEETING ABSTRACT; VISUALIZATION

METHOD

ORGANISM:

Organism Name

organisms (Organisms - Unspecified)

L3 ANSWER 16 OF 16 BIOSIS COPYRIGHT 2001 BIOSIS

ACCESSION NUMBER:

1992:156807 BIOSIS

DOCUMENT NUMBER:

BR42:73007

TITLE:

ATOMIC FORCE MICROSCOPY

IMAGING OF LARGE DOUBLE STRANDED DNA MOLECULES.
LYUBCHENKO Y L; GALL A A; SHLYAKHTENKO L S; HARRINGTON R E;

AUTHOR(S): LYUBCHENKO LINDSAY S M

CORPORATE SOURCE:

DEP. BIOCHEMISTRY, UNIV. NEVADA RENO, RENO, NV 89557.

SOURCE:

JOINT MEETING OF THE AMERICAN SOCIETY FOR BIOCHEMISTRY AND MOLECULAR BIOLOGY/BIOPHYSICAL SOCIETY, HOUSTON, TEXAS, USA, FEBRUARY 9-13, 1992. FASEB (FED AM SOC EXP BIOL) J, (1992.

6 (1), A149.

CODEN: FAJOEC. ISSN: 0892-6638.

DOCUMENT TYPE:

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FILE SEGMENT:

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LANGUAGE: CONCEPT CODE:

General Biology - Symposia, Transactions and Proceedings of

Conferences, Congresses, Review Annuals 00520 Genetics and Cytogenetics - General \*03502 Biochemical Studies - Nucleic Acids, Purines and

Pyrimidines \*10062

Biophysics - Molecular Properties and Macromolecules

\*10506

INDEX TERMS:

Miscellaneous Descriptors

ABSTRACT GENOMIC MAPPING NUCLEIC ACID-PROTEIN

COMPLEXES

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ANSWER 1 OF 18 CAPLUS COPYRIGHT 2001 ACS
                        2001:171629 CAPLUS
ACCESSION NUMBER:
                          Correction of: 1997:564938
DOCUMENT NUMBER:
                        134:178462
                          Correction of: 127:176339
                        Preparation of [(pyrrolidinoalkoxy)phenyl]benzothiophe
TITLE:
                        nes and analogs as thrombin inhibitors
                        Bastian, Jolie A.; Chirgadze, Nickolay Y.; Denney,
INVENTOR(S):
                        Michael L.; Foglesong, Robert J.; Harper, Richard W.;
                        Johnson, Mary G.; Klimkowski, Valentine J.; Kohn, Todd
                        J.; Lin, Ho-shen; Lynch, Michael P.;
                        Mccowan, Jefferson R.; Palkowitz, Alan D.; Richett,
                        Michael E.; Sall, Daniel J.; Smith, Gerald F.;
                        Takeuchi, Kumiko; Tinsley, Jennifer M.; Zhang,
                        Minsheng
                        Eli Lilly and Company, USA
PATENT ASSIGNEE(S):
                        PCT Int. Appl., 43 pp.
SOURCE:
                        CODEN: PIXXD2
DOCUMENT TYPE:
                        Patent
                        English
LANGUAGE:
INT. PATENT CLASSIF.:
                        A61K031-34
           MAIN:
                        A61K031-38; A61K031-40; A61K031-42; A61K031-395;
      SECONDARY:
                        A61K031-425; A61K031-435; A61K031-495; C07D211-06;
                        C07D241-02; C07D285-10; C07D307-78; C07D307-87;
                        C07D307-93; C07D333-52; C07D333-56; C07D333-66;
                        C07D333-72; C07D401-02; C07D403-14
CLASSIFICATION:
                        27-9 (Heterocyclic Compounds (One Hetero Atom))
                        Section cross-reference(s): 1
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
                                        APPLICATION NO. DATE
                    KIND DATE
    PATENT NO.
                     A1 19970717 WO 1996-US17995 19961030
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PRIORITY APPLN. INFO.:
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                                                                        P 19961009
                                                    US 1996-28252
                                                    WO 1996-US17995 W 19961030
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GRAPHIC IMAGE:

 $R^1$ 

R5

Ŗ4

Z

Т

ABSTRACT:

Title compds. [I; R1 = 1 or 2 of H, halo, Me, OMe, CONH2, etc.; R4 = 2122(CH2)j(CHR2)k(CH2)mNRaRb; R2 = OH, CH2OH, CO2Me; R5 = 232425(CH2)q(CHR3CHR3)rRcRd; R3 = H, alkyl, etc.; R3R3 = (CH2)3-4; Ra,Rb,Rc,Rd = H or alkyl; NRaRb,NRcRd = heterocyclyl; Z = O, S, CH:CH, CH2CH2; Z1,Z; 1,4-phenylene, (hetero)arylene; Z2 = bond, NH, CH2, O, S, NHCO; Z3 = O, S, CH:C, CO, C:CH2; Z5 = bond, NH, CH2, O, S, etc.; j,k,r = 0 or 1; m = 0-4; q = 0-2] were prepd. Thus, benzo[b]thiophene-2-boronic acid was condensed with 4-BrC6H4OMe and the product acylated by 4-(MeO)C6H4COCl to give, after hydrolysis, I [R1 = H, R4 = C6H4(OH)-4, R5 = COC6H4(OH)-4]. The latter was etherified by 1-(2-chloroethyl)pyrrolidine to give, after redn., I [R1 = H, R4 = R, R5 = CH2R, R = 4-(2-pyrrolidinoethyl)phenyl]. Data for biol. activity of 2 prepd. I were given.

SUPPL. TERM:

benzothiophene pyrrolidinoalkoxyphenyl prepn thrombin

inhibitor

INDEX TERM:

9002-04-4P, Thrombin

ROLE: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL

(Biological study); PREP (Preparation); USES (Uses)

(inhibitors; [(pyrrolidinoalkoxy)phenyl]benzothiophenes
and analogs)

INDEX TERM:

193960-53-1P 193960-54-2P 193960-55-3P 193960-56-4P 19396C-60-CP 193960-57-5P 193960-58-6P 193960-61**-**1P :9396C-64-4P 193960-65**-**5P 193960-62-2P 193960-63-3P 193960-67-72 193960-68-8F 193960-66-6P 193962-69-98 193960-71-3P 193960-72-4P 193960-73-5P 193960-70-2P 193960-74-6P 193960-75-7P 193960-76-8P 193960-78-0P 193960-81-5P 193960-82-6P 193960-79-1P 193960-80-4P 193960-84-8P 193960-85-9P 193960-86-0P 193960-83-7P 193960-90-6P 193960-92-8P 193960-87-1P 193960-88-2P 193960-95-1P 193960-97-3F 193960-98-4P 193960-94-0P 193960-99-5P 193961-01-2P 193961-03-4P 193961-05-6P 193961-09-0P 193961-11-4P 193961-12-5P 193961-07-8P 193961-16-9P 193961-13-6P 193961-15-8P 193961-17-0P 193961-23-8P 193961-19-2P 193961-21-6P 193961-25-0P 193961-29**-**4P 193961-30-7P 193961-31-8P 193961-27-2P 193961-37-4P 193961-33-0P 193961-35-2P 193961-39-6P 193961-45-4P 193961-43-2P 193961-47-6P 193961-41**-**0P 193961-51-2P 193961-53-4P 193961-55-6F 193961-49-8P 193961-59-0P 193961-61-4P 193961-63-6P 193961-57-8P 193961-67-0P 193961-69-2P 193961-71-6P 193961-65-8P 193961-75-0P 193961-77-2P 193961-79-4P 193961-73-8P 193961-87-4P 193961-83-0P 193961-85-22 193961-81**-**8P 193961-91-0P 193961-92-1P 193961-93-2P 193961-89-6P 193361-95-4F 193961-96-58 193961-94-3P 193961-99-8P 193962-00-4F 193962-01-5P 193961-98-7P 193962-03-7P 193962-04-8P 193962-05-9P 193962-02-6P 193962-07-1P 193962-08-2P 193962-09-3P 193962-06-0P 193962-11**-**7P 193962-13-9P 193962-14-0P 193962-10-6P 193962-17-3P 193962-18-4P 193962-19-5P 193962-15-1P 193962-22-0P 193962-20-8P 193962-21-9P 193962-23-1P 193962-25-3P 193962-26-4P 193962-27-5F 193962-24-2P 193962-29-7P 193962-30-0P 193962-31-1P 193962-28-6P 193962-33-3P 193962-34-4P 193962-35-5P 193962-32-2P 193962-36-6P 193962-37-7P 193962-38-8P 193962-39-9P 193962-41-3F 193962-42-4P 193962-43-5E 193962-40-2P 193962-**4**5-7P 193962-**4**6-8P 193962-48-0P 193962-44-6P 193962-50-4P 193962-51-5P 193962-52-6P 193962-49-1P 193962-56-0P 193962-58-2P 193962-60-6P 193962-54-8P 193962-63-9P 193962-65-1P 193962-67-3P 193962-62-8P 193962-72-0P 193962-71-9P 193962-69-5P 193962-70-8P 193962-77-5P 193962-74-2P 193962-75-3P 193962-73**-**1P

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ROLE: BAC (Biological activity or effector, except adverse);
SPN (Synthetic preparation); THU (Therapeutic use); BIOL
(Biological study); PREP (Preparation); USES (Uses)
   (prepn. of [(pyrrolidinoalkoxy)phenyl]benzothiophenes and
   analogs as thrombin inhibitors)
193964-12-4P
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ROLE: BAC (Biological activity or effector, except adverse);
SPN (Synthetic preparation); THU (Therapeutic use); BIOL
(Biological study); PREP (Preparation); USES (Uses)
   (prepn. of [(pyrrolidinoalkoxy)phenyl]benzothicphenes and
   analogs as thrombin inhibitors)
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                                          99-58-1,
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4-hydroxybenzoate 100-07-2, 4-Anisoyl chloride
                                                  100-35-6,
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110-91-8, Morpholine, reactions 122-04-3, 4-Nitrobenzoyl chloride 123-75-1, Pyrrolidine, reactions 150-76-5,
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619-58-9, 4-Iodobenzoic acid 624-28-2, 2,5-Dibromopyriding
696-62-8, 4-Iodoanisole 758-16-7, N,N-
Dimethylthioformamide 816-40-0, 1-Bromo-2-butanone
876-08-4, 4-Chloromethylbenzoyl chloride
                                         1007-16-5.
3-Bromo-4-fluorobenzoic acid 1081-73-8,
1-[2-(4-Bromophenoxy)] ethyl) pyrrolidine 1200-07-3,
4-Bromocinnamic acid 1461-25-2, Tetrabutyltin 2033-76-3,
1=(2-Bromoethoxy)-4-chlorobenzene 2176-98-9,
Tetrapropyltin 2362-12-1, 4-Bromo-2-methylphenol
2426-87-1, 4-Benzyloxy-3-methoxybenzaldehyde 2955-88-6,
1-Pyrrolidineethanol 3535-37-3, 3,4-Dimethoxybenzoyi
chloride 3556-83-0, Methyl 3-methoxy-4-methylben2cate
3647-69-6, 4- 2-Chloroethyl)morpholine hydrochloride
3943-74-6, Methyl vanillate 4397-53-9,
4-Benzyloxybenzaldehyde 4465-44-5, L-Serine,
N-triphenylmethyl-, methyl ester 4654-39-1,
4-Bromobenzeneethanol 5326-23-8, 6-Chloronicotinic acid
5335-87-5, Bis(4-methoxyphenyl) disulfide 6091-44-7,
Piperidine hydrochloride 6232-88-8, 4-Bromomethylbenzoic
      7217-59-6, 2-Methoxybenzenethiol 7250-67-1,
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INDEX TERM:

INDEX TERM:

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pyrrolidinyl)ethyl]benzoate
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ACCESSION NUMBER:
                        1999:625538 CAPLUS
DOCUMENT NUMBER:
                        131:351185
                        Solid phase chemistry approach to the SAR development
                        of a novel class of active site-directed thrombin
                         inhibitors
                         Johnson, Mary George; Bronson, Duane D.; Gillespie,
                        Jan E.; Gifford-Moore, Donetta S.; Kalter, Kyomi;
                        Lynch, Michael P.; McCowan, Jefferson R.;
                        Redick, Catherine C.; Sall, Daniel J.; Smith, Gerald
                         F.; Foglesong, Robert J.
CORPORATE SOURCE:
                        Sphinx Pharmaceuticals, A Division of Ell Lilly and
                        Company, Durham, NC, 27707, USA
                        Tetrahedron (1999), 55(39), 11641-11652
                        CODEN: TETRAB; ISSN: 0040-4020
                        Elsevier Science Ltd.
                        Journal
                        English
                        27-9 (Heterocyclic Compounds (One Hetero Atom))
                        Section cross-reference(s): 1
A solid phase chem. approach utilizing Mitsunobu chem., amine
functionalization, and parallel purifn. was used to produce a diverse library
of benzothiophene analogs. These analogs were used to advance the SAR of this
class of mols. and give new directions for future studies.
                  benzothiophene solid phase prepn thrombin inhibitor
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                  ROLE: BAC (Biological activity or effector, except adverse;
                  SPN (Synthetic preparation); BIOL (Biological study); PREP
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INDEX TERM:

TITLE:

AUTHOR(S):

SOURCE:

PUBLISHER:

LANGUAGE:

ABSTRACT:

SUPPL. TERM:

INDEX TERM:

INDEX TERM:

DOCUMENT TYPE:

CLASSIFICATION:

(solid phase prepn. of benzothiophenes as active site-directed thrombin inhibitors) 250651-25-3 250651-27-5 INDEX TERM: ROLE: RCT (Reactant) (solid phase prepn. of benzothiophenes as active site-directed thrombin inhibitors) INDEX TERM: 166975-45-7DP, polymer bound ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (solid phase prepn. of benzothiophenes as active site-directed thrombin inhibitors) INDEX TERM: 250651-26-4P ROLE: SPN (Synthetic preparation); PREP (Preparation) (solid phase prepn. of benzothiophenes as active site-directed thrombin inhibitors) REFERENCE COUNT: 24 REFERENCE(S): (1) Barnett, H; Hemostasis and Thrombosis: Basic Principles and Clinical Practice, 2nd ed 1987, P1301 (2) Brady, S; J Med Chem 1998, V41, P401 CAPLUS (3) Castro, J; J Org Chem 1994, V59, P2289 CASLUS (4) Colman, R: Hemostasis and Thrombosis: Basic Principles and Clinical Practice, 3rd ed 1994, P3 (5) Colman, R; Hemostasis and Thrombosis: Basic Principles and Clinical Practice, 3rd ed 1994, P508 (6) Edit, J; J Clin Invest 1989, V84, P18 (7) Edmunds, J; Ann Rep Med Chem 1996, V31, P51 CAPLUS (8) Farall, M; J Org Chem 1976, V41, P3877 (9) Heras, M; Circulation 1989, V79, P657 CAPLUS (10) Hirsh, J; Hemostasis and Thrombosis: Basic Principles and Clinical Practice, 2nd ed 1987, P1063 (11) Hirsh, J; Hemostasis and Thrombosis: Basic Principles and Clinical Practice, 2nd ed 1987, Pl199 (12) Hughes, D; Organic Preparations and Procedures Int 1996, V28, P127 CAPLUS (13) Lumma, W; J Med Chem 1998, V41, P1011 CAPLUS (14) Machovich, R; The Thrombin 1984, V1, P1 CAPLUS (15) Mann, K: Hemostasis and Thrombosis: Basic Principles and Clinical Practice, 3rd ed 1994, P184 (16) Sall, D; J Med Chem 1997, V4C, P3489 CAPLUS (17) Scarborough, R; Ann Rep Med Chem 1995, V3C, FTI CAPLUS (18) Smith, G; New Anticoagulants for the Cardiovascular Patient 1997, P265 (19) Tsunoda, T; Chem Lett 1994, P539 CAPLUS (20) Tsunoda, T; Tetrahedron Lett 1993, V34, P1639 CAPLUS (21) Tsunoda, T; Tetrahedron Lett 1994, V35, P5081 CAPLUS (22) Tsunoda, T; Tetrahedron Lett 1995, V36, P2529 CAPLUS (23) Tsunoda, T; Tetrahedron Lett 1995, V36, P2531 CAPLUS (24) Wiley, M; Exp Opin Ther Patents 1997, V7, Pl265 CAPLUS ANSWER 3 OF 18 CAPLUS COPYRIGHT 2001 ACS 1998:721694 CAPLUS ACCESSION NUMBER: 129:343408 DOCUMENT NUMBER: TITLE: Preparation of 3-benzyl(or benzoyl)-2phenylbenzo[b]thiophenes as antithrombotic agents Bach, Nicholas J.; Bastian, Jolie A.; Chirgadze, INVENTOR(S): Nickolay Y.; Denney, Michael L.; Foglesong, Robert J.; Harper, Richard W.; Johnson, Mary G.; Lin, Ho-shen; Lynch, Michael P.; McCowan, Jefferson R.; Palkowitz, David A.; Sall, Daniel J.; Smith, Gerald F.; Takeuchi, Kumiko; Zhang, Minsheng Eli Lilly and Co., USA PATENT ASSIGNEE(S): SOURCE: PCT Int. Appl., 138 pp. CODEN: PIXXD2

(Preparation)

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Patent
DOCUMENT TYPE:
                                       English
LANGUAGE:
INT. PATENT CLASSIF.:
                                       C07D409-14
                  MAIN:
                                       27-9 (Heterocyclic Compounds (One Hetero Atom))
CLASSIFICATION:
                                       Section cross-reference(s): 1, 63
FAMILY ACC. NUM. COUNT:
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       PATENT NO.
                                KIND DATE
                                                                   APPLICATION NO.
                                                                                              DATE
       WO 9849161
                                  A1
                                            19981105
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A1 20000223
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PRIORITY APPLN. INFO.:
                                                               US 1997-44297 19970430
                                                               WO 1998-US883C
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GRAPHIC IMAGE:
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            R6
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## ABSTRACT:

НО

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The title compds. [I; A = C(0), CH2; D = CH, C(0H), CMe, C(0Me), N; E = CH, CMe, C(0Me), C(halo); R2 = X2(CH2)mNRaRb (wherein X2 = a direct bond, CH2, O, S; m = 1-5; provided that when m = 1, then X2 = a direct bond; Ra, Rb = H, C1-3 alkyl; NRaRb = pyrrolidino, piperidino, morpholino); R3 = ON(Rg)2, SRh, CORi,

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etc. (wherein R9 = Me, Et; Rh = 2-thiazolyl; Ri = MeO); R6 = H, OH, MeO] and their pharmaceutically acceptable salts, useful as thrombin inhibitors, were prepd. and formulated. Thus, e.g., a multi-step synthesis of the title compd. II.oxalate, starting with 4-benzyloxybenzaldehyde, was described. Compds. I are effective at  $0.01-1000 \, \text{mg/kg/day}$ .

SUPPL. TERM: antithrombotic agent benzylphenylbenzothiophene benzoylphenylbenzothiophene prepn formulation; thrombin inhibitor benzylphenylbenzothiophene benzoylphenylbenzothiophene prepn formulation; benzothiophene prepn formulation antithrombotic INDEX TERM: Antithrombotics Thrombin inhibitors (prepn. of 3-benzyl(or benzoyl)-2phenylbenzo(b)thiophenes as antithrombotic agents; INDEX TERM: 215605-16-6P 215605-40-62 ROLE: BAC (Biological activity or effector, except adverse ; RCT (Reactant); SPN (Synthetic preparation); THO (Therapeutic use); BIOL (Biological study); PREF (Preparation); USES (Uses) (prepn. of 3-benzyl(or benzoyl)-2phenylbenzo[b]thiophenes as antithrombotic agents) INDEX TERM: 215604-83-4P 215604-84-5P 215604-86-7P 215604-88-9P 215604-91-4P 215604-89-0P 215604-90-3P 215604-92-5P 215604-93-6P 215604-94-7P 215604-95-8P 215604-96-9P 215604-97-0P 215604-98-1P 215604-99-2P 215605-00-8P 215605-07-5P 215605-02-0P 215605-04-2P 215605-06-4P 215605-08-6P 215605-C9-7P 215605-10-0P 215605-11-19 215605-14-4P 215605-12-2P 215605-13-3P 215605-15-5P 215605-19-9P 215605-17-7P 215605-18-8P 215605-20-2P 215605-22-4P 215605-23-5P 215605-24-6P 215605-25-7P 215605-26-8P 215605-27-9P 215605-29-1P 215605-28-0P 215605-30-4P 215605-31-5P 215605-32-6P 215605-33-7P 215605-34-8P 215605-35-92 215605-37-1P 215605-39-3P 215605-41-7P 215605-43-92 2156C5-44-0P ROLE: BAC (Biological activity or effector, except adverse ; SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREF 'Preparation'; USES (prepn. of 3-benzyl or benzoyl)-2phenylbenzo[b]thiophenes as antithrombotic agents) INDEX TERM: 85-41-6, Phthalimide 105-36-2, Ethyl bromoacetate 120-92-3, Cyclopentanone 121-34-6, Vanillic acid 286-20-4, 7-Oxabicyclo[4.1.0]heptane 288-94-8, 1H-Tetrazole 403-16-7, 3-Chloro-4-fluorobenzoic acid 403-43-0, p-Fluorobenzoyl chloride 497-25-6, 2-0xazolidone 766-00-7, 2-Cyclopentylethanol 1081-73-8 2417-72-3, Methyl 4-bromomethylbenzoate 2749-11-3, (S)-2-Amino-1-propanol 2969-81-5, Ethyl 4-bromobutyrate 3195-95-7, 2-Pyrrolidone sodium salt 3445-11-2, 1-(2-Hydroxyethyl)-2-pyrrolidinone 3556-83-0, Methyl 3-methoxy-4-methylbenzoate 3699-54-5, 1-(2-Hydroxyethyl)-2-imidazolidinone 3710-84-7, N,N-Diethylhydroxylamine 4397-53-9, 4-Benzyloxybenzaldehyde 5454-83-1, Methyl 5-bromovalerate 5685-05-2, 2-Mercaptothiazole 7368-78-7, 4-Bromoguaiacol 7697-28-1, 4-Bromo-3methylbenzoic acid 13865-19-5, Methyl 3-formylpropionate 18190-44-8, N-(2-Hydroxyethyl)succinimide 25542-62-5, 34743-88-9 30321-23-3 Ethyl 6-bromohexanoate 77470-83-2, 4-Chloromethyl-2-58029-83-7 63675-96-7 methylthiazole hydrochloride 90560-10-4, 6-Methoxybenzo[b]thiophene 104901-43-1, Methyl 3-bromo-4-methylbenzoate 193964-19-1 193966-49-3

193966-83-5

ROLE: RCT (Reactant)

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(prepn. of 3-benzyl(or benzoyl)-2-
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ACCESSION NUMBER: 1998:719268 CAPLUS
DOCUMENT NUMBER:
                            129:343406
TITLE:
                            Preparation of benzothiophenes as antithrombotic
                            agents
INVENTOR(S):
                            Chirgadze, Nickolay Y.; Denney, Michael L.; Fisher,
                            Matthew J.; Foglesong, Robert J.; Harper, Richard W.;
                            Johnson, Mary G.; Lin, Ho-shen; Lynch, Michael
                            P.; McCowan, Jefferson R.; Miller, Shawn C.;
                            Palkowitz, Alan D.; Richett, Michael E.; Sall, Danier
                            Jon; Smith, Gerald Floyd; Takeuchi, Kumiko; Zhang,
                            Minsheng
PATENT ASSIGNEE(S):
                            Eli Lilly and Co., USA
SOURCE:
                            PCT Int. Appl., 171 pp.
                            CODEN: PIXXD2
DOCUMENT TYPE:
                            Patent
LANGUAGE:
                            English
INT. PATENT CLASSIF.:
             MAIN:
                            A61K031-445
        SECONDARY:
                            A61K031-38
CLASSIFICATION:
                            27-9 (Heterocyclic Compounds (One Hetero Atom,)
                            Section cross-reference(s): 1, 63
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
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     WO 9848804
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                        Al 19981105
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EP 1998-922073

19980430

Al 19981124

20000816

A1

AU 9874700

EP 1027051

R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, PT, IE

PRIORITY APPLN. INFO.:

US 1997-45188 P 19970430

WO 1998-US8717 W 19980430

OTHER SOURCE(S): GRAPHIC IMAGE:

MARPAT 129:343406

\* STRUCTURE DIAGRAM TOO LARGE FOR DISPLAY - AVAILABLE VIA OFFLINE PRINT \*

#### ABSTRACT:

The title compds. [I; A = C(O), CH2; D = CH, CMe, C(OMe), N; E = CH, CMe, C(OMe), C(halo), N; R2 = NRaCO(CH2)mRb, OCH2Rb, NHCORg, etc. (wherein m = 0-1; Ra = H, Me; Rb = II, III (G = O, S, NH, CH2, CH2CH2; Rc = H, Me; L = NH, NMe, CH2; Rg = 5-membered heteroarom. having 2 heteroatoms selected from O, S and  $\kappa$ in which the carbonyl group is bonded to a ring carbon situated between a ring heteroatom and another ring carbon, etc.)); R3 = X3(CH2)sNRsRt (X3 = a direct bond, CH2, O; s = 1-2; provided that when s = 1, then X3 = a direct bond; and Rs and Rt = H, C1-3 alkyl or NRsRt = pyrrolidino, piperidino, morpholino); R6 -H, OH, MeO] and their pharmaceutically acceptable salts, useful as thrombin inhibitors, were prepd. and formulated. Thus, a multi-step synthesis of the title compd. (R)-IV oxalate, starting with 6-methoxybenzo[b]thiophene, is described. Compds. I are effective at 0.01-1000 mg/kg/day.

SUPPL. TERM:

benzothiophene prepn formulation antithrombotic; thrombin

inhibitor benzothiophene preph formulation

INDEX TERM:

Antithrompotics Thrombin inhibitors

(prepn. of benzothiophenes as antithrombotic agents)

215387-83-0P INDEX TERM:

ROLE: BAC (Biological activity or effector, except adverse);

RCT (Reactant); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP

(Preparation); USES (Uses)

(prepn. of benzothiophenes as antithrombotic agents)

INDEX TERM:

215386-99-5P 215387-00-1P 215387-02-3P 215387-03-4P 215387-04-5P 215387-C5-6P 215387-08-9P 215387-09-0P 215387-10-3P 215387-11-4P 215387-13-6P 215387-14-7P 215387-16-9P 215387-17-0P 215387-15-8P 215387-18-1P 215387-20-5P 215387-21-6P 215387-19~2P 215387-23-8P 215387-24-9P 215387-25-CP 215387-26-1P 215387-28-3P 215387-34-1P 215387-30-7P 215387-33-0P 215387-29-4P 215387-38-5P 215387-39-6P 215387-40-9P 215387-36-3P 215387-41-0P 215387-43-22 215387-44-3P 215387-45-4P 215387-46-5P 215387-48-7P 215387-49-8F 215387-51-2P 215387-55-6P 215387-52-3P 215387-54-5P 215387-56-7P 216387-60-3P 215387-59-0P 215387-57-8P 215387-59-9P 215387-65-8P 215387-63-6P 215387-64-7P 215387-62-5P 215387-70-5P 215387-66-92 215387-68-1P 215387-69-2P 215387-71-6P 215387-72-7P 215387-73-8P 215387-74-9P 215387-75-0P 215387~76**-**1P 215387-77-2P 215387-78-3P 215387-79-4P 215387-82-9P 215387-80-7P 215387-81-82 215387-89-6P 215387-85-2P 215387-87-4P 215387-84-1P 215387-92-1P 215387-93-2P 215387-94-3P 215387-91-0P 215387-96-5P 215387-98-7P 215387-95-4P 215388-00-4P 215388-04-8P 215388-01-5P 215388-02-6P 215388-03-7P 215388-06-0P 215388-08-2P 215388-10-68 215388-05-9P 215388-18-4P 215388-13-9P 215388-17-3P 215388-12-8P ROLE: BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses)

INDEX TERM:

(prepn. of benzothiophenes as antithrombotic agents) 55-22-1, Isonicotinic acid, reactions 65-85-0, Benzoic

acid, reactions 79-44-7, Dimethylcarbamoyl chloride 96-30-0, 2-Chloro-N-methylacetamide 98-09-9, Benzenesulfonyl chloride 98-78-2 98-79-3, 2-Pyrrolidinone-5-carboxylic acid 104-92-7, 4-Bromoaniscle 106-41-2, 4-Bromophenol 122-04-3, 4-Nitrobenzoyl chloride 123-75-1, Pyrrolidine, reactions 403-43-0, 4-Fluorobenzoyl 534-03-2 586-78-7, 1-Bromo-4-mitrobenzene chloride 611-73-4, Benzoylformic acid 623-00-7, 4-Bromobenzonitrile 628-20-6, 4-Chlorobutyronitrile 766-00-7, 2-Cyclopentylethanol 1072-84-0, 4-Imidazolecarboxylic acid 1192-63-8, 1-Pyrrolidinecarbonyl chloride 1621-91-6, 3-Pyrazolecarboxylic acid 2362-12-1 2675-89-0, 2-Chloro-N, N-dimethylacetamide 2955-88-6, 1-(2-Hydroxyethyl)pyrrolidine 3209-71-0. 3-Isoxazolecarboxylic acid 3445-11-2, 1-(2-Hydroxyethyl)-2pyrrolidinone 3556-83-0, Methyl 3-methoxy-4-methylbenzoate 3699-54-5, 1-(2-Hydroxyethyl)-2-imidazolidinone 3770-22-7,6-Oxo-2-piperidinecarboxylic acid 3943-74-6, Methyl 4-hydroxy-3-methoxybenzoate 4042-36-8, D-Pyroglutamic and 4397-53-9, 4-Benzyloxybenzaldehyde 5089-33-8, 4-Bromo-N, N-bis(trimethylsilyl)aniline 5332-06-9, 4-Bromobutyronitrile 7250-67-1, 1-(2-Chloroethyl)pyrrolidine hydrochloride 7268-43-1 15159-40-7, 4-Morpholinecarbonyl chloride 17342-08-4, (S)-5-Hydroxymethyl-2-pyrrolidinone 19771-63-2 21169-71-1, 5-Isoxazolecarboxylic acid 21277-16-7 24856-58-4, p-Bromobenzaldehyde dimethylacetal 26690-80-2 32741-98-3 34743-88-9, 2-(4-Bromophenoxy)ethanol 42346-68-9 35320-23-1 40584-55-2 52574-06-8 54120-42-2, 3-Bromo-N, N-bis(trimethylsilyl)aniline 63675-90-1 63675-91-2 64520-53-2 66673-40-3 73918-56-6 76453-38-8 80522-42-5, Triisopropylsilyl 90560-10-4, trifluoromethanesulfonate 84449-68-3 6-Methoxybenzo[b]thiophene 97899-36-0 98437-23-1, Benzo[b]thiophene-2-boronic acid 194901-43-1, Methyl 3-bromo-4-methylbenzoate 175460-94-3, 6-Methoxy-2-(4hydroxyphenyl)benzo[b]thicphene 193964-97-5 193964-98-6 215386-71-3 215386-76-8 215388-71-9 215386-65-5 ROLE: RCT (Reactant) (prepn. of benzothiophenes as antithrombotic agents) 15546-08-4P 27884-09-9P 63676-06-2P 63676-12-0P 70264-94-7P 78946-25-5P 106391-86-0P 65540-08-1P 182133-35-3P 193964-66-8P 193964-75-9P 193964-77-1P 193965-55-8P 193965-93-4F 193965-54-7P 193965-94-5P 193966-03-9P 193966-02-8P 193966-04-0P 193966-06-2P 193966-35-7P 193966-77-7P, 193966-34-6P 1-Bromo-4-(triisopropylsilyloxy)benzene 193966-78-8P 193966-79-9P 193966-80-2P 215378-95-3P 215378-96-4P 215379-15-0P 215379-57-0P 215379-00-3P 215379-09-2P 215380-13-5P 215380-08-8P 215382-00-6P 215382-22-2P 215382-37-9P 215382-97-1P 215382-32-4P 215382-64-2P 215383-02-1P 215383-70-3P 215383-72-5P 215383-91-8P 215383-94-1P 215383-99-6P 215384-02-4P 215384-17-1P 215384-46-6P 215384-57**-**9P 215384-62-6P 215384-28-4P 215384-91-1P 215387-32-9P 215388-21-9P 215388-22-0P 215388-24-2P 215388-23-1P 215388-26-4P 215388-27-5P 215388-28-6P 215388-29-7P 215388-30-0F 215388-31-1P 215388-33-3P 215388-32-2P 215388-35-5P 215388-37-7P 215388-39-9F 215388-40-2P 215388-38-8P 215388-41-38 215388-42-4P 215388-43-5P 215388-44-6P 215388-45-7F 215388-46-8P 215388-47-99 215388-49-1F 215388-50-4P 215388-51-5P 215388-52-6P 215388-53-7P 215388-54-8P 215388-58-2P 215388-55-9P 215388-57-1P 215388-59-3P 215388-61-7P 215388-63-9P 215388-64-0P 215388-62-8P

INDEX TERM:

215388-67-3P 215388-68-4P 215388-66-2P 215388-69-5P

215388-70-8P

ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. of benzothiophenes as antithrombotic agents)

ANSWER 5 OF 18 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1998:719264 CAPLUS

DOCUMENT NUMBER:

130:3845

TITLE:

Preparation of 1-benzyl-2-phenylbenzimidazoles as

antithrombotic agents

INVENTOR(S):

Bastian, Jolie A.; Chirgadze, Nickolay Y.; Denney, Michael L.; Fisher, Matthew J.; Foglesong, Robert J.; Harper, Richard W.; Johnson, Mary G.; Klimkowski,

Valentine J.; Lin, Ho-shen; Lynch, Michael P.

; McCowan, Jefferson R.; Miller, Shawn C.; Mullaney, Jeffrey T.; Richett, Michael E.; Sall, Damiel J.; Smith, Gerald F.; Takeuchi, Kumiko; Tinsley, Jennifer M.; Wiley, Michael R.; Zhang, Minsheng

PATENT ASSIGNEE(S):

SOURCE:

Eli Lilly and Co., USA PCT Int. Appl., 75 pp.

CODEN: PIXXD2

DOCUMENT TYPE:

LANGUAGE:

Patent English

INT. PATENT CLASSIF.:

MAIN:

A61K031-415

SECONDARY:

A61K031-41

CLASSIFICATION:

28-9 (Heteropyclic Compounds (More Than One Hetero

Atom.))

Section cross-reference(s): 1, 63

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO. KIND DATE APPLICATION NO. DATE A1 199811C5 WO 1998-US8755 1998C430 WO 9848800 W: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, GM, GW, HU, ID, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, SD, SZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG EP 1998-920057 19980430 SB, GR. TT 1-AU 1998-72707 AU 9872707 A1 19981124 EP 1019047 A1 20000719 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, NL, SE, PT, IE, FI APPLN. INFO.:

US 1997-45335 P 19970501

WO 1998-US8755 W 19980430

URCE(S):

MARPAT 130-3845 PRIORITY APPLN. INFO.:

OTHER SOURCE(S): MARPAT 130:3845

GRAPHIC IMAGE:

Rl

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 $R^2$ 

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OMe

N N

N

Ν

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O N

HO N

ΙI

### ABSTRACT:

The title compds. [I; E = CH, CMe, C(OMe), C(halo); R denotes 0-2 substituents selected from halo, Me, Et, etc.; R1 = 5-tetrazolylmethyl, 2-carboxypyrrolidinelylmethyl, etc. R2 = 2-(1-pyrazolyl) ethoxy, 2-(1-pyrrolidinyl) ethoxy, etc.] and their pharmaceutically acceptable salts, useful as thrombin inhibitors, were prepd. and formulated. Thus, a multi-step synthesis of the title compd. II as its dioxalate salt, starting with 4-amino-3-nitrophenol, was described. Preferred compds. I reduce the net clot wt. to approx. 25-30% of control, or even lower, at an i.v. dose of 33.176 .mu.mol/kg/h.

SUPPL. TERM:

benzylphenylbenzimidazole prepn formulation antithrombotic;

thrombin inhibitor benzylphenylbenzimidazole prepn

formulation

INDEX TERM:

Antithrombotics
Thrombin inhibitors

(prepn. of 1-benzyl-2-phenylbenzimidazoles as

antithrombotic agents)

INDEX TERM:

215656-86-3P 215656-88-5P 215656-90-9P 215656-91-0F

215656-92-1P 215656-93-2P 215656-94-3P 215656-96-5P

215656-97-6P 215656-98-7P

ROLE: BAC (Biological activity or effector, except adverse);

SPN (Synthetic preparation); THU (Therapeutic use); BIOL

(Biological study); PREP (Preparation); USES (Uses)

(prepn. of 1-benzyl-2-phenylbenzimidazoles as antithrombotic agents)

INDEX TERM:

50-85-1, 4-Methylsalicylic acid 98-88-4, Benzoyi chloride

99-96-7, reactions 100-46-9, Benzylamine, reactions

288-13-1, Pyrazole 610-81-1, 4-Amino-3-nitrophenol 616-45-5, 2-Pyrrolidinone 1493-27-2, 2-Fluoronitrobenzene

5587-42-8, Imidazole sodium sait 7250-67-1 = 40958-42-1, Pyrazole sodium salt 41253-21-8, 1,2,4-Triazole sodium

salt 63675-91-2 69731-93-7 193964-75-9

ROLE: RCT (Reactant)

(prepn. of 1-benzyl-2-phenylbenzimidazoles as

antithrombotic agents)

INDEX TERM:

99-76-3P 56850-91-0P 63675-89-8P 63675-90-1P

74733-27-0P 81245-24-1P 122893-33-8P 193964-76-0P 215656-53-4P 215656-54-5P 215656-55-6P 215656-60-3P 215656-61-4P 215656-64-7P 215656-65-8P 215656-66-9P 215657-02-6P 215657-03-7P 215657-04-8P 215657-05-9P 215657-06-0P 215657-00-0P 215657-ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation) (prepn. of 1-benzyl-2-phenylbenzimidazoles as antithrombotic agents)

REFERENCE COUNT:

REFERENCE(S):

(1) Lunn; US 5552426 A 1996 CAPLUS (2) Narr; US 5541229 A 1996 CAPLUS

ANSWER 6 OF 18 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1993:124535 CAPLUS

DOCUMENT NUMBER:

118:124535

TITLE:

Preparation of phenylimidazolone herbicides and

intermediates

INVENTOR(S):

Crouse, Gary D.; Lynch, Michael P.; Webster,

Jeffery D.; Wright, John P.

PATENT ASSIGNEE(S):

USA

SOURCE:

U.S., 36 pp.

CODEN: USXXAM

DOCUMENT TYPE:

Patent English

LANGUAGE: INT. PATENT CLASSIF.:

MAIN:

C07D211-40

SECONDARY:

C07D267-12

US PATENT CLASSIF.:

540451000

CLASSIFICATION:

28-9 (Heterocyclic Compounds (More Than One Hetero

Atom))

Section cross-reference(s): 5

FAMILY ACC. NUM. COUNT: 1

PATENT INFORMATION:

PATENT NO.				KIND		DATE		APPLICATION NO.					υ.	DATE			
									-	<del>-</del> -							
US	5153	316		А		1992	1006		U	S 19	91-7	1022	3	1991	0604		
WO	WO 9221662			A1		19921210			WO 1992-US4716					19920604			
	W:	AT,	ΑU,	BB,	BG,	BR,	CA,	CH,	DE,	DΚ,	ES,	FI,	GB,	HU,	JP,	ΚP,	KR,
		LK,	LU,	MG,	MW,	NL,	NO,	PL,	RO,	RU,	SD,	SE,	US				
	RW:	AT,	BE,	BF,	ВJ,	CF,	CG,	CH,	CI,	CM,	DE,	DK,	ES,	FR,	GA,	GB,	GN,
		GR,	ΙT,	LU,	MC,	ML,	MR,	NL,	SE,	SN,	TD,	TG					
AU 9221623 A1 1						19930108 AU 1992-21623							19920604				
PRIORITY APPLN. INFO.:								US 1991-710223					19910604				
								;	#C 1	a 92 - 1	US47:	ř.		1992	2674		

OTHER SCURCE(S):

MARPAT 118:124535

GRAPHIC IMAGE:

### ABSTRACT:

Herbicides [I; A = bond, C:T; R = H, alkyl, haloalkyl, cyanoalkyl, Ph; Rl = H, halo, (halo)alkyl, cyano; when A = bond, RRl = atoms to form an (unsatd.) (substituted) 3-7 membered carbocyclic ring; T = O, S; U = H; V = OH; or CV = bond; W = halo, cyano; when W = halo, then Rl = cyano; X = H, halo; Y = H, halo, cyano, alkyl, CH3, OCF3; Z = H, halo, OH, alkyl, aryloxy, allyl, (acyl)amino, NO2, alkoxy; YZ = atoms to form a (substituted) (unsatd.) 3-7 membered carbocyclic ring], and intermediates II (n = 1-4, other variables as above), were prepd. Thus, 1-(4-chlorophenyl)-4-formyl-5-chloro-3-propyl-1,3-dihydro-2H-imidazol-2-one was stirred with NH2OH.HCl in EtOH/H2O and the product was stirred with Cl3CCOCl/Et3N in CH2Cl2 to give title compd. III. III at 0.5 lb/acre preemergent gave complete control of; jimsonweed, foxtail millet, lambsquarter, etc.

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phenylimidazolone prepn herbicide; imidazolone phenyl
SUPPL. TERM:
                   herbicide
INDEX TERM:
                   Herbicides
                      (phenylimidazolones)
INDEX TERM:
                   145861-66-1P
                                  145861-67-2P
                                                  145861-68-3P
                                                                 145861-69-4P
                   145861-70-7P
                                   145861-71-8P
                                                  145861-72-9P
                                                                 145861-73-0P
                   145861-74-1P
                                   145861-75-2P
                                                  145861-76-3P
                                                                  145861-77-4P
                   145861-78-5P
                                   145861-79-6P
                                                  145861-80-9P
                                                                  145861-81-0P
                   145861-82-1P
                                   145861-83-2P
                                                  145861-84-3P
                                                                 145861-85-4P
                   145861-86-5P
                                   145861-87-6P
                                                  145861-88-7P
                                                                 145861-89-8P
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                                                  145861-92-3P
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                                                                  145861-97-8P
                   145861-94-5P
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                                                  145861-96-7P
                   145861-98-9P
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                                                  145862-00-6P
                                                                 145862-01-7F
                   145862-02-82
                                   145862-03-9P
                                                  145862-04-0P
                                                                 145862-05-1P
                                                                 145862-09-5P
                   145862-06-2P
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                   145862-10-8P
                                   145862-11-9P
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                                   145862-19-7P
                                                  145862-20-0P
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                   145862-30-2P
                                   145862-31-3P
                                                  145862-32-4P
                                                                  145862-33-5P
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                                   145862-35-7P
                                                  145962-36-8P
                                   145862-39-12
                                                                 145862-41-5F
                   145862-38-0P
                                                  145862-40-4F
                   145862-42-6P
                                  145862-43-7P
                                                  145862-44-8P
                   ROLE: AGR (Agricultural use); BAC (Biological activity or
                   effector, except adverse); SPN (Synthetic preparation); BIOL
                   (Biological study); PREP (Preparation); USES (Uses)
                      (prepn. of, as herbicide)
INDEX TERM:
                   6330-25-2P
                                70928-90-4P 145862-50-6P
                                                            145862-51-7P
                   145862-52-8P
                                  145862-53-9P 145862-54+0P 145862-55-1P
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145862-56-2P 145862-57-3P 145862-58-4P 145862-59-5P 145862-61-9P 145862-60-8P 145862-62-0P 145862-63-1P 145862-66-4P 145862-64-2P 145862-65-3P 145962-67-5F 145862-71-1P 145862-70-0P 145862-68-6P 145862-69-7P 145862-72-2P 145862-73-3P 145862-74-45 149862-15-51 ROLE: SPN (Synthetic preparation ; PREP (Preparation (prepn. of, as herbicide intermediate) INDEX TERM: 75-44-5, Phosgene 104-12-1, 4-Chlorophenylisocyanate 105-56-6, Ethyl cyanoacetate 105-60-2, .epsilon.-Caprolactam, reactions 106-47-8, p-Chloroaniline, reactions 624-83-9, Methyl isocyanate 2216-92-4, Phenylglycine ethyl ester 2521-89-3 91167-85-0 108310-38-9 84478-65-9 145862-45-9 145862-46-0 145862-47-1 145862-48-2 145862-49-3 ROLE: RCT (Reactant) (reaction of, in prepn. of herbicide) 74-89-5, Methylamine, reactions 107-97-1, Sarcosine 675-20-7, .delta.-Valerolactam 5470-11-1, Hydroxylamine INDEX TERM: hydrochloride 25808-30-4 32315-10-9, Triphosgene 70591-20-7 145862-76-6 ROLE: RCT (Reactant) (reaction of, in prepn. of phenylimidazolone herbicide) INDEX TERM: 75-45-6, Chlorodifluoromethane 106-96-7, Propargyl bromide 107-30-2, Chloromethyl methyl ether 616-45-5. 2-Pyrrolidinone 18668-72-9 ROLE: RCT (Reactant) (reaction of, in preph. of pheylimidazolone herbloide, ANSWER 7 OF 18 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1993:22158 CAPLUS DOCUMENT NUMBER: 118:22158 TITLE: Synthesis and gametocidal activity of 1-aryl-5-(aminocarbonyl)-1H-pyrazole-4-carboxylic AUTHOR(S): Lynch, Michael P.; Ackmann, Stephen A.; Heim, Dale R.; Davis, George E.; Staszak, Michael A.; Beck, James R.; Tschabold, Edward E.; Wright, Fred L. DowElanco Res. Lab., Greenfield, IN, 46140, USA CORPORATE SOURCE: SOURCE: ACS Symp. Ser. (1992), 504 (Synth. Chem. Agrochem. III), 200-11 CODEN: ACSMC8; ISSN: 0097-6156 DOCUMENT TYPE: Journal; General Review LANGUAGE: English CLASSIFICATION: 28-0 (Heterocyclic Compounds (More Than One Hetero Atom)) Section cross-reference(s): 5 ABSTRACT: A series of 1-aryl-5-(aminocarbonyl)-1H-pyrazole-4-carboxylic acids were serendipitously discovered to be chem. hybridizing agents. Different synthetic routes were developed for the active analogs depending on whether an electron withdrawing group or electron donating group was present on the Ph ring. Development of the "second generation gametocides" produced analogs which were 5-6 times more active than the original lead. A review with 20 refs. SUPPL. TERM: review gametocide prepn aminocarbonyl pyrazciecarboxylate; hybridization aminocarbonyl yrazolecarboxylate prepn review INDEX TERM: Genetics (hybridization agents, (aminocarbonyl)pyrazolecarboxylate

INDEX TERM: 145147-06-4D, 5-(Aminocarbonyl)-lH-pyrazole-4-carboxylic acid, 1-aryl derivs.

INDEX TERM:

s)

Plant hormones and regulators

(gametocides, (aminocarbonyl)pyrazolecarboxylates)

ROLE: RCT (Reactant)

ROLE: RCT (Reactant)

(hybridization agents and gametocides)

ANSWER 8 OF 18 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1991:122156 CAPLUS

DOCUMENT NUMBER:

114:122156

TITLE:

1-Alkyl-5-cyano-1H-pyrazole-4-carboxamides. Synthesis

and herbicidal activity

AUTHOR(S):

Lynch, Michael P.; Beck, James R.; Tao,

Eddie V. P.; Aikins, James; Babbitt, George E.; Rizzo,

John R.; Waldrep, T. William

CORPORATE SOURCE:

Lilly Res. Lab., Eli Lilly and Co., Greenfield, IN,

46140, USA

SOURCE:

ACS Symp. Ser. (1991), 443(Synth. Chem. Agrochem. 2),

144-57

CODEN: ACSMO8; ISSN: 0097-6156

DOCUMENT TYPE:

Journal

LANGUAGE:

English

CLASSIFICATION:

28-8 (Heterocyclic Compounds (More Than One Hetero

Atom))

Section cross-reference(s): 5

GRAPHIC IMAGE:

Me<sub>3</sub>C CN

N

Ν

CONHMe

### ABSTRACT:

A symposium on the investigation of EL-177, 5-cyano-1-(1,1-dimethylethyl.-Nmethyl-1H-pyrazole-4-carboxamide, (I) as a new and effective preemergent corn and postemergent cereal herbicide is described. A variety of 1-alkyl-5-cyano-1H-pyrazole-4-carboxamides were prepd. regioselectively using tertiary carbocation chem. With olefins incapable of forming tertiary carbocations, a direct method of alkylating pyrazoles under basic conditions was examd. Regioisomers produced using this method were sepd. by chromatog. Identification of the regioisomers was made by an empirical method comparing the solvent shifts of the pyrazole proton in DMSO-D6 and CDC13. A comparison of the herbicidal activity of the various pyrazole carboxamides is presented.

SUPPL. TERM:

alkylcyanopyrazolecarboxamide prepn herbicide activity symposium; EL177 alkylcyanopyrazolecarboxamide prepn

herbicide activity symposium; pyrazolecarboxamide

cyanodimethylethylmethyl prepn herbicide activity symposium; structure activity alkylcyanopyrazolecarboxamide herbicide

symposium

INDEX TERM:

Herbicides

INDEX TERM:

(alkyloyanopyrazolecarboxamides) Molecular structure-biological activity relationship (herbicidal, of alkyloyanopyrazolecarboxamides)

INDEX TERM:

98477-C7-7P

ROLE: BAC (Biological activity or effector, except adverse;; SPN (Synthetic preparation); BIOL (Biological study); PREF

(Preparation)

(prepn. and herbicidal activity of)

ANSWER 9 OF 18 CAPLUS COPYRIGHT 2001 ACS 1.1

ACCESSION NUMBER: 1990:405585 CAPLUS

DOCUMENT NUMBER: 113:5585

TITLE: Carbon-13 NMR chemical shifts of 1-alkyl-3(5)-cyano-18-

pyrazole-4-darboxylic atid esters

AUTHOR(S): Babbitt, George E.; Lynch, Michael P.; Beck,

James R.

CORPORATE SOURCE: Lilly Res. Lab., Eli Lilly and Co., Greenfield, IN,

46140, USA

SOURCE: Magn. Reson. Chem. (1990), 28(1), 90-2

CODEN: MRCHEG; ISSN: 0749-1581

DOCUMENT TYPE: Journal

LANGUAGE: English

CLASSIFICATION: 22-10 (Physical Organic Chemistry)

ABSTRACT:

A compilation of 13C NMR chem. shifts for 13 pairs of 3- and 5-cyano-substituted pyrazole regioisomers is reported. All of the ring carbon and cyano carbon 13C chem. shifts show a regular, predictable correlation with the particular isomer, whether 3-cyano or 5-cyano. These shifts occurred in very narrow ranges, precluding any confusion of assignment within the group of compds. studied. X-ray crystallog. anal. was performed on one of the samples.

SUPPL. TERM: NMR alkylcyanopyrazolecarboxylate ester; pyrazolecarboxylate

ester alkyloyano NMR carbon

INDEX TERM: Nuclear magnetic resonance

(of alkyloyanopyrazolebarboxylate esters, darbon=13)

INDEX TERM: 33090-55-0 33090-56-1 98477-12-4 119741-58-1

119741-59-2 119741-60-5 121485-81-2 121909-85-1 121909-87-3 121909-86-2 121909-88-4 121909-89-5 121909-91-9 121909-90-8 121909-92-0 127526-68-5 127526-70-9 127526-71-0 127526-69-6 127526-72-1 127526-74-3 127526-73-2 127526-75-4 127526-76-5

127526-77-6 127553-52-0 ROLE: PRP (Properties) (carbon-13 NMR of)

INDEX TERM:

14762-74-4

ROLE: PRP (Properties)

(nuclear magnetic resonance, of

alkylcyanopyrazolecarboxylate esters, carbon-13)

L1 ANSWER 10 OF 18 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER: 1990:114072 CAPLUS

DOCUMENT NUMBER: 112:114072

TITLE: Synthesis and herbicidal activity of 1-aryl-5-halo and

1-aryl-5-(trifluoromethyl)-1H-pyrazole-4-carboxamides

AUTHOR(S): Waldrep, Thomas W.; Beck, James R.; Lynch,

Michael P.; Wright, Fred L.

CORPORATE SOURCE: Lilly Res. Lab., Greenfield, IN, 46140, USA

SOURCE: J. Agric. Food Chem. (1990), 38(2), 541-4

CODEN: JAFCAU; ISSN: 0021-8561

DOCUMENT TYPE: Journal LANGUAGE: English

CLASSIFICATION: 5-3 (Agrochemical Bioregulators)
Section cross-reference(s): 28

GRAPHIC IMAGE:

 $R^2 R^1 R$ 

CONR4R5

R3 N

N I

ABSTRACT:

A series of 1-aryl-5-halo- and 1-aryl-5-(trifluoromethyl)-1H-pyrazole-4-carboxamides (I, R = Cl, CF3; R1 = R2 = R3 = H, Cl, CF3, CMe, Me; R4 = cyclopropyl, H, Et, Bu, etc.; R5 = Et, Me, OMe) exhibit moderate to strong herbicidal activity in preemergence and postemergence tests. At 1/2 lb/acre, corn, rice, wheat, cotton, and soybean show tolerance, while large crabgrass, foxtail millet, common lambsquarters, redroot pigweed, wild mustard, velvetleaf, jimsonweed, and zinnia were killed or severely injured. A total of 83 5-halo analogs and 47 5-trifluoromethyl analogs were synthes:zed and their herbicidal activities detd. to examine the structure-activity relationships. The order of activity at C-5 of the pyrazole ring was CF3 > Cl .simeq. Br > I. The order of activity involving substitution on the carboxamide molety was cyclopropyl .simeq. Me > di-Me > Et > iso-Fr. Substitution on the benzene ring did not result in any major increase in activity when compared with the corresponding Ph analog.

SUPPL. TERM: herbicide halopyrazolecarboxamide deriv INDEX TERM: Herbicides (arylhalo and arylfluoromethylpyrazolecarboxamides) INDEX TERM: Molecular structure-biological activity relationship (herbicidal, of arylhalo and arylfluoromethylpyrazolecarboxamides) INDEX TERM: 125024-12-6 ROLE: BIOL (Biological study) (fluoromethylpyrazolecarboxamide herbicides prepn. from) INDEX TERM: 98477-04-4P 98533-16-5P 98533-18-7P 98533-21-2P 98533-32-5P 98533-29-0P 98533-31-4P 98533-33**-**62 98533-36-9P 98533-44-9P 98533-48-3P 98533-49-42 98533-56-3P 98533-57-4P 98533-53-0P 98533-55-28 98533-59-6P 98533-60-9P 98533-65-4P 98533-71-2P 98533-75-6P 98533-79-0P 98533-83-6P 98533-85-8P 98533-97-0P 98533-89-2P 98533-90-5P 98533-93-8P 98533-95-0P 98533-97-2P 98533-98-3P 98533-99-48 98534-00-02 98534-01-1P 98534-02-2P 98534-04-4P 98534-22-6P 98534-06-62 98534-23-7P 98534-24-8P 98534-27-1P 98534-25**-**9P 98534-26-0P 98534-28-2P 98534-31-7P 98534-29-3P 98534-30-6P 98534-32-8P 98534-36-2P 98534-33-9P 98534-34-0P 98534-35-1P 98534-37-3P 98534-38-4P 98534-39-5P 98534-40-8P 98534-41-9P 98534-43-1P 98534-44-2P 98534-45-3P 98534-47-5P 98534-48-6P 98534-49-7P 98534-50-0P 98534-52-2P 98534-53-3P 98534-54-4P 98534-55-5P 98534-57-7P 98534-58-8P 98534-59-9P 98534-56-6P 98534-61-3P 102996-38-3P 98534-60-2P 125024-11-5P 125048-87-5P ROLE: AGR (Agricultural use); BAC (Biological activity or

L1 ANSWER 11 OF 18 CAPLUS COPYRIGHT 2001 ACS

to)

ACCESSION NUMBER:

1989:477904 CAPLES

DOCUMENT NUMBER:

111:77904

TITLE:

Alkylation studies with 5-cyano-IH-pyranole-4-

effector, except adverse); SPN (Synthetic preparation); BIOI

(prepn. and herbicidal activity of, structure in relation

(Biological study); PREP (Preparation); USES (Uses)

carboxylic acid, ethyl ester

AUTHOR(S):

Beck, James R.; Aikins, James; Lynch, Michael

P.; Rizzo, John R.; Tao, Eddie V. P.

CORPORATE SOURCE:

Lilly Res. Lab., Div. Eli Lilly and Co., Greenfield,

IN, 46140, USA

SOURCE:

J. Heterocycl. Chem. (1989), 26(1), 3-6

CODEN: JHTCAD; ISSN: 0022-152X

DOCUMENT TYPE:

Journal

LANGUAGE:

English

CLASSIFICATION:

28-8 (Heterocyclic Compounds (More Than One Hetero

Atom) '

Section cross-reference(s): 5

OTHER SCURCE(S): GRAPHIC IMAGE:

CASREACT 111:77904

CN

CN

CN

HN

PrMe2CN

Me 3CN N

N

CO2Et I

CO2Et II

CONHMe 111

ABSTRACT:

The title compd. (I) was alkylated regioselectively at N(1) by tertiary carbocations utilizing H2SO4 catalysis and relatively mild conditions. Thus, I was treated with Me2C:CHMe and H2SO4 to give 64% cyano(dimethylpropyl)pyrazolec arboxylate II. In the presence of BF3, the alkylation occurred regionelectively at N(2). Reaction of I with alkyl halides under basic conditions resulted in mixts, of the two isomers with alkylation at  $N(\theta)$ predominating.

SUPPL. TERM:

regioselective alkylation cyanopyrazolecarboxylate;

alkylcyanopyrazolecarboxamide prepn herbicide;

alkylcyanopyrazolecarboxylate

INDEX TERM:

Herbicides

(cyanomethylpyrazolecarboxamides)

INDEX TERM:

Regiochemistry

(of alkylation of cyanopyrazolecarboxylate with alkenes

and alkyl halides)

INDEX TERM:

Alkenes, reactions ROLE: RCT (Reactant)

(regioselective alkylation by, by

cyanopyrazolecarboxylate;

INDEX TERM:

Alkylation

(regioselective, of cyanopyrazolecarboxylate with alkenes

and alkyl halides)

INDEX TERM:

33090-56-1P 33090-55-0P 121909-88-4P 121909-89-5P

121909-91-9P 121909-92-0P 121909-90-8P

ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and amidation of, with methylamine)

INDEX TERM:

119741-58-1P

ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and amidation of, with methylamines;

INDEX TERM:

119741-56-9P

ROLE: BAC (Biological activity or effector, except adverse.; SPN (Synthetic preparation); BIOL (Biological study); PREP

(Preparation)

(prepn. and herbicidal activity of)

INDEX TERM:

119741-59-2P 119741-60-5P 121909-85-1P 121909-86-2P

121909-87-3P

ROLE: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

INDEX TERM:

137-43-9, Cyclopentyl bromide 108-85-0, Cyclohexyl bromide

513-35-9, 2-Methyl-2-butene 693-89-0 760-21-4

763-29-1, 2-Methyl-1-pentene 816-79-5, 3-Ethyl-2-pentene

ROLE: RCT (Reactant)

(regioselective alkylation by, of

cyanopyrazolcarboxylate)

INDEX TERM:

119741-57-0

ROLE: RCT (Reactant)

(regioselective alkylation of, with alkenes and alkyl

ANSWER 12 OF 18 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1989:212704 CAPLUS

DOCUMENT NUMBER:

110:212704

TITLE:

Synthesis of ethyl 2-[(l-aryl-1H-1,2,4-triazo]-3-

yl)oxy]propionates and related derivatives Beck, James R.; Babbitt, George E.; Lynch,

Michael P.

CORPORATE SOURCE:

Lilly Res. Lab., Eli Lilly and Co., Greenfield, IN,

46140, USA

SOURCE:

J. Heterocycl. Chem. (1988), 25(5), 1467-70

CODEN: JHTCAD; ISSN: 0022-152X

DOCUMENT TYPE:

Journal

LANGUAGE:

AUTHOR(S):

English

CLASSIFICATION:

28-10 (Heterocyclic Compounds (More Than One Hetero

Atom))

OTHER SOURCE(S): GRAPHIC IMAGE:

CASREACT 110:212704

RN N

OH I N

# ABSTRACT:

Alkylation of 1-aryl-1H-1,2,4-triazol-3-cls I (R = Ph, 2-, 3-, 4-ClC6H5, 3-CF3C6H4, 2,4-Cl2C6H4, 3,4-Cl2C6H4) with MeCHBrCO2Et under basic conditions resulted in the formation of 2-[(1-aryl-1H-1,2,4-triazol-3-yl)oxy]propionic acid, Et esters. No N-alkylated products were detected. Similar alkylation of 2-oxo-5-phenyl-1,3,4-thiadiazole and the corresponding 1,3,4-oxadiazole gave only N-alkylated derivs. With 4-hydroxy-6-phenylpyrimidine and 2-oxo-4-phenylthiazole, both O- and N-alkylation occurred. assignments were based on IR and 13C-NMR spectral data.

SUPPL. TERM:

aryltriazolyloxypropionate; aryltriazolol alkylation

bromopropionate; triazolol aryl alkylation bromopropionate; thiadiazole oxophenyl alkylation bromopropionate; oxadiazole

oxophenyl alkylation bromopropionate; pyrimidine

hydroxyphenyl alkylation bromopropionate; thiszule excehengi

alkylation bromopropionate

INDEX TERM:

Regiochemistry

(of alkylation of triazolols, oxothiazoles,

oxooxadiazoles, oxothiadiazoles, and hydroxypyrimidine by

bromopropionate;

INDEX TERM:

Ring closure and formation

(of arylsemicarbazide with orthoformate, aryltriazolol

from)

INDEX TERM:

Alkylation

(of triazoles, oxothiazoles, oxocxadiazoles, oxothiadiazoles, and hydroxypyrimidines by

bromopropionate)

INDEX TERM:

Heterocyclic compounds

ROLE: SPN (Synthetic preparation); PREP (Preparation'

```
arylsemicarbazide with orthoformate;
                                                        24028-40-8
                   1199-02-6
                               3884-31-9
                                           4891-69-4
INDEX TERM:
                   ROLE: RCT (Reactant)
                       (alkylation of, with bromopropionate)
                   103-03-7 14577-00-5
                                           14580-27-9
                                                        14580-28-0
                                                                     14657-26-2
INDEX TERM:
                   42158-58-7
                                57802-85-4
                   ROLE: RCT (Reactant)
                      (cyclization of, with orthoformate, aryltriazolol from)
                   7727-37-9P
INDEX TERM:
                   ROLE: SPN (Synthetic preparation); PREP (Preparation)
                       (heterocyclic compounds, nitrogen, aryltriazolols, prepn.
                      of, via cyclization of arylsemicarbazide with
                      orthoformate)
                                37176-51-5P
INDEX TERM:
                   4231-68-9P
                                              42158-59-8P
                                                             84456-06-4P
                                 110626-11-4P
                   84456-12-2P
                   ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP
                   (Preparation)
                       (prepn. and alkylation of, with bromopropionate)
INDEX TERM:
                   23875-84-5P
                   ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP
                   (Preparation)
                       (prepn. and alkylation of, with bromopropionate or
                      bromoacetate)
INDEX TERM:
                   120590-04-7P
                   ROLE: SPN (Synthetic preparation); PREP (Preparation)
                       (prepn. of)
                                                                  11000/-58-41
INDEX TERM:
                   110607-55-1P
                                  110607-56-2P
                                                  110607-57-3F
                                                  110607-65-3P
                                  110607-6C-8P
                                                                 120590-05-8₽
                   110607-59-5P
                                  120590-07-0P
                                                  120590-08-1P
                                                                 120590-09-22
                   120590-06-9P
                   120590-10-5P
                   ROLE: AGR (Agricultural use); BAC (Biological activity or
                   effector, except adverse); SPN (Synthetic preparation.; BIOL
                   (Biological study); PREP (Preparation); USES (Uses)
                      (prepn. of, as plant growth regulator)
    ANSWER 13 OF 18 CAPLUS COPYRIGHT 2001 ACS
                         1989:8100 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         110:8100
TITLE:
                         Synthesis of 1-aryl-1H-pyrazolecarbonitriles and
                         related derivatives
AUTHOR(S):
                         Beck, James R.; Lynch, Michael P.; Wright,
                         Fred L.
                         Lilly Res. Lab., Eli Lilly and Co., Greenfield, IN,
CORPORATE SOURCE:
                         46140, USA
                         J. Heterocycl. Chem. (1988), 25(2), 555-8
SOURCE:
                         CODEN: JHTCAD; ISSN: 0022-152X
DOCUMENT TYPE:
                         Journal
LANGUAGE:
                         English
CLASSIFICATION:
                         28-8 (Heterocyclic Compounds (More Than One Hetero
                         Atom;)
                         CASREACT 110:8100
OTHER SOURCE(S):
GRAPHIC IMAGE:
                                R^3
        CN
                                     R^2
             CO2Et
R1C6H4N
                        R1C6H4N
                                         ΙI
                    Ι
```

(nitrogen, aryltriazolols, prepn. of, via cyclization of

ABSTRACT: Chloropyrazoles were treated with NaCN to give nitriles I (RI = H, Cl). I were converted to pyrazoles II (R2 = CO2H, CONH2, cyano, H, NH2, CO2Me; R3 = CONH2, CO2H, CO2Me, CONHMe). SUPPL. TERM: pyrazolecarbonitrile carboxy; cyanopyrazolecarboxylic acid INDEX TERM: 7664-41-7, Ammonia, reactions ROLE: RCT (Reactant) (amidation by, of pyrazolecarboxylic acid deriv.) INDEX TERM: 74-89-5, Methylamine, reactions ROLE: RCT (Reactant) (amidation by, of pyrzolecarbonyl chloride deriv.) 117766-97-9P INDEX TERM: ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and Curtius reaction of) INDEX TERM: 117778-62-8P ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and amidation of) INDEX TERM: 103053-10-7P ROLE: SPN (Synthetic preparation); PREP (Preparation) (prepn. and conversion of, to dicacarboxamide) INDEX TERM: 117767-01-8P ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and esterification of, by methanol) INDEX TERM: 98476-09-6P 98476-16-5P ROLE: SPN (Synthetic preparation); PREP (Preparation) (prepn. and hydration of) INDEX TERM: 117766-90-2P 117766-99-1P ROLE: RCT (Reactant); SPN (Synthetic preparation); FREP (Preparation) (prepn. and hydrolysis of) INDEX TERM: 117766-91-3P ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reaction of, with thionyl chloride) INDEX TERM: 98477-01-1P 103053-08-3P 117766-92-4P ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP (Preparation) (prepn. and reactions of) INDEX TERM: 117767-C2-9P ROLE: RCT (Reactant); SPN (Synthetic preparation); PREF (Preparation) (prepn. and selective sapon. of) INDEX TERM: 103053-21-0P 117766-93-5P 117766-94-6P 117766-95-7P 117766-96-8P 117767-00-7P 117767-03-0P 117767-C4-1P ROLE: SPN (Synthetic preparation); PREP (Preparation) (prepn. of) INDEX TERM: 117766-98-0 ROLE: RCT (Reactant) (prepn. reaction of, with sodium azide) INDEX TERM: 143-33-9, Sodium dyanide ROLE: RCT (Reactant) (substitution reaction of, with chloropyrazolecarboxylate esters) INDEX TERM: 98534-74-8 98534-76-0 ROLE: RCT (Reactant)

(substitution reaction of, with sodium cyanide)

L1 ANSWER 14 OF 18 CAPLUS COPYRIGHT 2001 ACS ACCESSION NUMBER: 1988:75280 CAPLUS

DOCUMENT NUMBER: 108:75280

TITLE: Synthesis of 1-(1,1-dimethylethyl)-1H-pyrazole-4-

carboxylate ester derivatives

Book lames R : Tynch Michael F

AUTHOR(S): Beck, James R.; Lynch, Michael P.

CORPORATE SOURCE: Lilly Res. Lab., Div. Eli Lilly and Co., Greenfield,

IN, 46140, USA

SOURCE: J. Heterocycl. Chem. (1987), 24(3), 693-5

CODEN: JHTCAD; ISSN: 0022-152X

DOCUMENT TYPE: Journal

LANGUAGE: English

CLASSIFICATION: 28-8 (Heterocyclic Compounds (More Than One Hetero

Atom))

Section cross-reference(s): 75

OTHER SOURCE(S): CASREACT 108:75280

GRAPHIC IMAGE:

CN R1

Me3CN CONHMe Me3CN CO2Et

N I N II

ABSTRACT:

Pyrazolecarboxamide deriv. I was prepd. from pyrazolecarboxylate ester deriv. II (R1 = Me) via II (R1 = CH2Br) and II (R1 = CH0). The cyclocondensation reaction of Me3CNHNH2 with MeCOC(:CHNMe2)CO2Et gave II (R1 = Me).

SUPPL. TERM: cyanopyrazolecarboxamide prepn crystal structure;

pyrazolecarboxamide cyano; pyrazolecarboxylate ester

INDEX TERM: Crystal structure

Molecular structure

(of cyanopyrazolecarboxamide deriv.)

INDEX TERM: 74-89-5, reactions

ROLE: RCT (Reactant)

(amidation by, of pyrazolecarboxylate ester deriv.)

INDEX TERM: 141-97-9

ROLE: RCT (Reactant)

(condensation reaction of, with DMF acetal)

INDEX TERM: 4637-24-5, DMF dimethyl acetal

ROLE: RCT (Reactant)

(condensation reaction of, with acetoacetate ester)

INDEX TERM: 94-05-3, Ethyl cyano(ethoxymethylene)acetate

ROLE: RCT (Reactant)

(cycloaddn.-cyclocondensation reaction of, with

alkylhydrazine)

INDEX TERM: 7400-27-3

ROLE: RCT (Reactant)

(cyclocondensation reaction of, with

(aminomethylene)acetoacetate ester deriv.)

INDEX TERM: 98477-12-4P

ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and amidation of, by methylamine)

INDEX TERM: 98477-08-8P

ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and bromination of)

INDEX TERM: 98477-09-9P

ROLE: SPN (Synthetic preparation); PREP (Preparation)

(prepn. and conversion of, to formylpryazolecarboxylate

analog)

INDEX TERM:

51145-57-4P

ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and cyclocondensation reaction of, with

alkylhydrazine)

INDEX TERM:

112779-11-0P 112779-12-1P

ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and dehydration of)

INDEX TERM:

ROLE: RCT (Reactant); SPN (Synthetic preparation); FREP

(Preparation)

(preph. and oximation of)

INDEX TERM:

112779-14-3P

98477-10-2P

ROLE: RCT (Reactant); SPN (Synthetic preparation); FREP

(Preparation)

(prepn. and reaction with nitrosyl chloride and hydrogen.

chloride)

INDEX TERM:

98477-07-7P 112779-13-2P

ROLE: SPN (Synthetic preparation); PREP (Preparation)

(prepn. of)

ANSWER 15 OF 18 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1987:575941 CAPLUS

DOCUMENT NUMBER:

107:175941

TITLE:

Nonaqueous diazotization of 5-amino-1-aryl-1H-pyrazole-

4-carboxylate esters

AUTHOR(S):

Beck, James R.; Gajewski, Robert P.; Lynch,

Michael P.; Wright, Fred L.

CORPORATE SOURCE:

Lilly Res. Lab., Eli Lilly and Co., Greenfield, IN,

46140, USA

SOURCE:

J. Heterocycl. Chem. (1987), 24(1), 267-70

CODEN: JATCAD; 188N: 00/2-152X

DOCUMENT TYPE:

LANGUAGE:

Journal English

CLASSIFICATION:

28-9 (Heterocyclic Compounds (More Than One Hetero

Atom))

OTHER SOURCE(S):

CASREACT 107:175941

ABSTRACT:

5-Amino-1-aryl-1H-pyrazole-4-carboxylate esters are converted to the corresponding desamino, chloro, bromo, icdo, and methylthic analogs by processes involving nonaq, diazotization. Diazotizing agents are alkyi nitrites except in the case of chlorine where nitrosyl chloride is used.

SUPPL. TERM:

diazotization aminopyrazolecarboxylate ester;

pyrazolecarboxylate ester; halopyrazolecarboxylate ester

INDEX TERM:

Diazotization

(of aminopyrazolecarboxylate esters, in prepn. of deamino

and halo analogs)

INDEX TERM:

3107-33-3 19690-59-6

ROLE: RCT (Reactant)

(cycloaddn.-cyclocondensation of, with cyano(ethoxymethylene)acetate ester)

INDEX TERM:

94-05-3, Ethyl cyano(ethoxymethylene)acetate

ROLE: RCT (Reactant)

(cycloaddn.-cyclocondensation of, with phenylhydrazines.

INDEX TERM:

15001-09-9

ROLE: RCT (Reactant)

(diazotization and reaction of, with bromine,

pyrazoloindazole deriv. from)

INDEX TERM:

15001-12-4

ROLE: RCT (Reactant)

(diazotization of, in prepn. of bromo analog)

INDEX TERM:

15001-13-5 16459-35-1

```
(diazotization of, in preph. of deamino and unloro
                       analogs)
INDEX TERM:
                   14678-87-6
                                 15001-08-8 16078-71-0
                   ROLE: RCT (Reactant)
                       (diazotization of, in prepn. of deamino and halo analogs)
INDEX TERM:
                   110821-47-1P
                   ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP
                    (Preparation)
                       (prepn. and cyclization of)
INDEX TERM:
                   110821-30-2P
                   ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP
                    (Preparation)
                       (prepn. and diazotization of, in prepn. of brominated
                       deamino compd.)
INDEX TERM:
                   110821-29-9P
                   ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP
                    (Preparation)
                       (prepn. and diazotization of, in prepn. of deaming and
                       halo analogs)
                                 .
1397-55-6P 98534-71-5P 98534-72-6P
98534-76-0P 103053-42-5P 110821-31-3P
INDEX TERM:
                   885-94-9P 91397-55-6P
                   98534-74-8P
                                                 110821-34-6F
110821-38-0P
                   110821-32-4P
                                  110821-33-5P
                                                                  110821-35-1P
                                                                  110821-39-18
                   110821-36-8P
                                   110821-37-9P
                                   110821-41-5P
                   110821-40-4P
                                                  110821-42-6P
                                                                 110821-43-7P
                                 110821-45-92
                                                                 110821-48-2P
                   110821-44-8P
                                                  110821-46-0P
                   ROLE: SPN (Synthetic preparation); PREP (Preparation)
                       (prepn. of)
INDEX TERM:
                   624-92-0, Dimethyl disulfide
                   ROLE: RCT (Reactant)
                       (reaction of, with pyrazolediazonium compd.)
     ANSWER 16 OF 18 CAPLUS COPYRIGHT 2001 ACS
                         1986:442791 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                         105:42791
TITLE:
                         Herbicidal 5-halo-1-halophenyl-1H-pyrazole-4-
                         carbonitriles
INVENTOR(S):
                         Beck, James R.; Lynch, Michael P.
PATENT ASSIGNEE(S):
                         Lilly, Eli, and Co. , USA
                         U.S., 10 pp. Cont.-in-part of U.S. Ser. No. 549,138,
SOURCE:
                         abandoned.
                         CODEN: USXXAM
DOCUMENT TYPE:
                         Patent
LANGUAGE:
                         English
INT. PATENT CLASSIF.:
            MAIN:
                         A01N043-56
       SECONDARY:
                         C07D231-14; C07D231-16
US PATENT CLASSIF.:
                         071092000
CLASSIFICATION:
                         28-8 (Heterocyclic Compounds (More Than One Hetero
                         Atom))
                         Section cross-reference(s): 5
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
     PATENT NO.
                    KIND DATE
                                           APPLICATION NO. DATE
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     US 4563210
                            19860107
                                                             19840913
                                           US 1984-650135
PRIORITY APPLN. INFO.:
                                        US 1983-549138
                                                             19831107
GRAPHIC IMAGE:
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ROLE: RCT (Reactant)

CN

Ν  $R_{\mathbf{n}}$ Ν Ι

ABSTRACT:

Pyrazolecarbonitriles I (n = 1, 2; R = F, C1; R1 = halo, CF3), which were prepd., exhibited herbicidal activity. A mixt. of EtOCH:C(CN;2, 2,3,4-Cl3C6H2NHNH2 (prepd. from the resp. aniline), HOAc, and water was refluxed to yield I (Rn = 2.3,4-Ci3, R1 = NH2), and the latter was treated with NOCl in CHCl3 to give I (Rn = 2, 3, 4-Cl3, R1 = Cl).

SUPPL. TERM: halopyrazolecarbonitrile prepn herbicide;

pyrazolecarbonitrile halo prepn herbicide

INDEX TERM: Herbicides

(halo(halophenyl)pyrazolecarbonitriles)

7664-41-7, reactions INDEX TERM: ROLE: RCT (Reactant)

(amidation by, of pyrazolecarboxylic acid deriv.)

INDEX TERM: 5446-18-4

ROLE: RCT (Reactant)

(cycloaddn.-cyclocondensation of, with (ethoxymethylene)acetoacetate ester deriv.)

INDEX TERM: 123-06-8 571-55-1

ROLE: RCT (Reactant)

(cycloaddn.-cyclocondensation of, with phenylhydrazine

deriv.)

INDEX TERM: 5879:-79-0

ROLE: RCT (Reactant)

(deamination-bromination of)

INDEX TERM: 79002-96-3 102996-25-8

ROLE: RCT (Reactant)

(deamination-chlorination of)

INDEX TERM: 98534-78-2P

ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and amidation of)

INDEX TERM: 80025-74-7P

ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and cycloaddn.-cyclocondensation of, with

(ethoxymethylene)malononitrile)

INDEX TERM: 80025-46-3P

ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and deamination-chlorination of)

INDEX TERM: 102996-38-3P

ROLE: RCT (Reactant); SPN (Synthetic preparation); PREP

(Preparation)

(prepn. and dehydration of)

INDEX TERM: 98534-77-1P

ROLE: RCT (Reactant); SPN (Synthetic preparation); PREF

(Preparation)

(prepn. and sapon. of)

INDEX TERM: 102996-24-7P 102996-23-6P 102996-26-92 102996-27-0F

> 102996-28-1P 102996-29-2P 102996-30-5P :02996-31-6F 102996-32-7P 102996-33-8P 102996-34-9P 102996-35-0P 102996-36-1P 102996-37-2P 102996-39-4P 102996-40-7P

ROLE: AGR (Agricultural use); BAC (Biological activity or effector, except adverse); SPN (Synthetic preparation); BIOI (Biological study); PREF (Preparation); USES (Pses)

(prepn. of, as herbicide)

INDEX TERM:

634-67-3

ROLE: RCT (Reactant)

(N-nitrosation of, and redn. of product from)

L1 ANSWER 17 OF 18 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1986:222875 CAPLUS

DOCUMENT NUMBER:

104:222875

TITLE:

Pyrophosphohydrolase activity and inorganic pyrophosphate content of cultured human skin

fibroblasts. Elevated levels in some patients with calcium pyrophosphate dihydrate deposition disease

AUTHOR(S):

Ryan, Lawrence M.; Wortmann, Robert L.; Karas, Barbara; Lynch, Michael P.; McCarty, Daniel

J.

CORPORATE SOURCE:

Dep. Med., Med. Coll. Wisconsin, Milwaukee, WI, 53226,

USA

SOURCE:

J. Clin. Invest. (1986), 77(5), 1689-93

CODEN: JCINAO; ISSN: 0021-9738

DOCUMENT TYPE:

Journal

LANGUAGE:

English

CLASSIFICATION:

14-11 (Mammalian Pathological Biochemistry)

ABSTRACT:

In Ca pyrophosphate dihydrate (CPPD) crystal deposition disease, metabolic abnormalities favoring extracellular inorg, pyrophosphate (PPi) accumulation have been suspected. Elevations of intracellular PPi in cultured skin fibroblasts from a single French kindred with familial CPPD deposition and elevated nucleoside triphosphate pyrophosphohydrolase activity (NTPPPH), which generates PPI in exts. of CPPD crystal-contg. cartilages favor this suspicion. To det. whether NTPPPH activity or PPi content of cells might be a disease marker expressed in extraarticular cells, human skin-derived fibroblasts were obtained from control donors and patients affected with the sporadic and familial varieties of CPPD (CPPD-S and CPPD-F) deposition. Intracellular PPi was elevated in both CPPD-S and CPPD-F fibroblasts compared with control fibroblasts. Ecto-NTPPPH activity was elevated in CPPD-S but not in CPPD-F. Intracellular PPi correlated with ecto-NTPPPH. Elevated PPi levels in skin fibroblasts may serve as a biochem. marker for patients with familial or sporadic CPPD crystal deposition disease; ecto-NTPPPH activity further separates the sporadic and familial disease types. Expression of these biochem. abnormalities in nonarticular cells implies a generalized metabolic abnormality.

SUPPL. TERM:

calcium pyrophosphate dinydrate deposition firmtlast

biochem; pyrophosphohydrolase calcium pyrophosphate

dehydrate deposition

INDEX TERM:

Fibroblast

(pyrophosphates and pyrophosphohydrolase of numan, in calcium pyrophosphate dihydrate crystal deposition

disease)

INDEX TERM:

17031-92-4

ROLE: BIOL (Biological study)

(crystals, deposition of, pyrophosphates and pyrophosphohydrolase of human fibroblasts in)

INDEX TERM:

9027-73-0 9033-44-7 9075-54-1

ROLE: BIOL (Biological study)

(of fibroblast, in calcium pyrophosphate dihydrate
crystal deposition disease in humans)

L1 ANSWER 18 OF 18 CAPLUS COPYRIGHT 2001 ACS

ACCESSION NUMBER:

1983:420607 CAPLUS

DOCUMENT NUMBER:

99:20607

TITLE:

Inorganic pyrophosphate levels in blood platelets from normal donors and patients with calcium pyrophosphate